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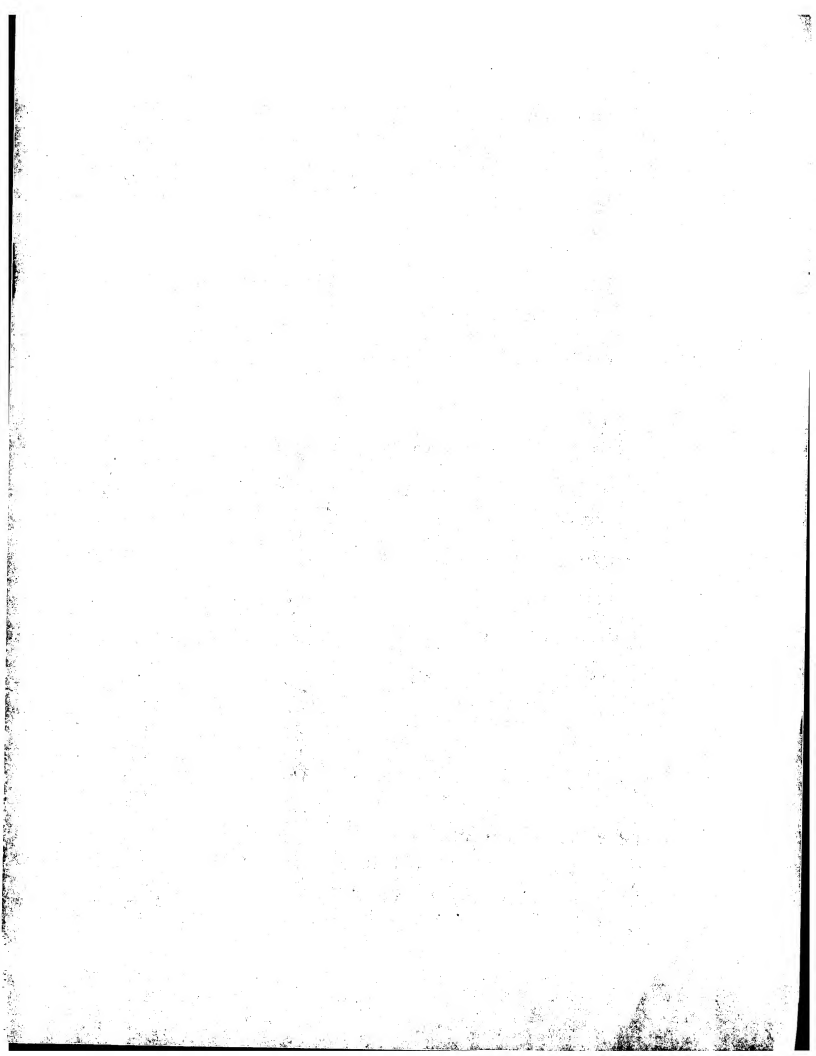
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### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classificati n <sup>6</sup>:

C12N 15/40, C07K 14/18, C12Q 1/70, C07K 16/10, A61K 39/29 (11) International Publication Number:

WO 99/04008

A2

(43) Internati nal Publication Date:

28 January 1999 (28.01.99)

(21) International Application Number:

PCT/US98/14688

(22) International Filing Date:

16 July 1998 (16.07.98)

(30) Priority Data:

60/053,062 09/014,416 18 July 1997 (18.07.97) 27 January 1998 (27.01.98)

US US

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Published

Without international search report and to be republished upon receipt of that report.

(54) Title: CLONED GENOMES OF INFECTIOUS HEPATITIS C VIRUSES AND USES THEREOF

#### (57) Abstract

The present invention discloses nucleic acid sequences which encode infectious hepatitis C viruses and the use of these sequences, and polypeptides encoded by all or part of these sequences, in the development of vaccines and diagnostics for HCV and in the development of screening assays for the identification of antiviral agents for HCV.

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Title Of Invention

Cloned Genomes Of Infectious Hepatitis C Viruses And Uses Thereof

This application claims the benefit of U.S. Provisional Application No. 60/053,062 filed July 18, 1997.

### Field Of Invention

The present invention relates to molecular approaches to the production of nucleic acid sequences which comprise the genome of infectious hepatitis C viruses. In particular, the invention provides nucleic acid sequences which comprise the genomes of infectious hepatitis C viruses of genotype la and lb strains. The invention therefore relates to the use of these sequences, and polypeptides encoded by all or part of these sequences, in the development of vaccines and diagnostic assays for HCV and in the development of screening assays for the identification of antiviral agents for HCV.

#### Background Of Invention

Hepatitis C virus (HCV) has a positive-sense single-strand RNA genome and is a member of the virus family Flaviviridae (Choo et al., 1991; Rice, 1996). As for all positive-stranded RNA viruses, the genome of HCV

for all positive-stranded RNA viruses, the genome of hove functions as mRNA from which all viral proteins necessary for propagation are translated.

The viral genome of HCV is approximately 9600 nucleotides (nts) and consists of a highly conserved 5' untranslated region (UTR), a single long open reading

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frame (ORF) of approximately 9,000 nts and a complex 3'
UTR. The 5' UTR contains an internal ribosomal entry site
(Tsukiyama-Kohara et al., 1992; Honda et al., 1996). The
3' UTR consists of a short variable region, a

polypyrimidine tract of variable length and, at the 3' end, a highly conserved region of approximately 100 nts (Kolykhalov et al., 1996; Tanaka et al., 1995; Tanaka et al., 1996; Yamada et al., 1996). The last 46 nucleotides

of this conserved region were predicted to form a stable stem-loop structure thought to be critical for viral replication (Blight and Rice, 1997; Ito and Lai, 1997; Tsuchihara et al., 1997). The ORF encodes a large

polypeptide precursor that is cleaved into at least 10 proteins by host and viral proteinases (Rice, 1996). The predicted envelope proteins contain several conserved N-linked glycosylation sites and cysteine residues (Okamoto et al., 1992a). The NS3 gene encodes a serine protease

and an RNA helicase and the NS5B gene encodes an RNA-dependent RNA polymerase.

Globally, six major HCV genotypes (genotypes 1-6) and multiple subtypes (a, b, c, etc.) have been identified (Bukh et al., 1993; Simmonds et al., 1993). The most divergent HCV isolates differ from each other by more than 30% over the entire genome (Okamoto et al., 1992a) and HCV circulates in an infected individual as a quasispecies of closely related genomes (Bukh et al., 1995; Farci et al., 1997).

At present, more than 80% of individuals infected with HCV become chronically infected and these chronically infected individuals have a relatively high

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risk of developing chronic hepatitis, liver cirrhosis and hepatocellular carcinoma (Hoofnagle, 1997). In the U.S., HCV genotypes la and 1b constitute the majority of infections while in many other areas, especially in Europe and Japan, genotype 1b predominates.

The only effective therapy for chronic hepatitis C, interferon (IFN), induces a sustained response in less than 25% of treated patients (Fried and Hoofnagle, 1995). Consequently, HCV is currently the most common cause of end stage liver failure and the reason for about 30% of liver transplants performed in the U.S. (Hoofnagle, 1997). In addition, a number of recent studies suggested that the severity of liver disease and the outcome of therapy may be genotype-dependent (reviewed in Bukh et al., 1997). In particular, these studies suggested that infection with HCV genotype 1b was associated with more severe liver disease (Brechot, 1997) and a poorer response to IFN therapy (Fried and Hoofnagle, 1995). As a result of the inability to develop a universally effective therapy against HCV infection, it is estimated that there are still more than 25,000 new infections yearly in the U.S. (Alter 1997) Moreover, since there is no vaccine for HCV, HCV remains a serious public health problem.

However, despite the intense interest in the development of vaccines and therapies for HCV, progress has been hindered by the absence of a useful cell culture system and the lack of any small animal model for laboratory study. For example, while replication of HCV in several cell lines has been reported, such observations have turned out not to be highly reproducible. In

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addition, the chimpanzee is the only animal model, other than man, for this disease. Consequently, HCV has been able to be studied only by using clinical materials obtained from patients or experimentally infected chimpanzees (an animal model whose availability is very limited).

However, several researchers have recently reported the construction of infectious cDNA clones of HCV, the identification of which would permit a more effective search for susceptible cell lines and facilitate molecular analysis of the viral genes and their function. For example, Dash et al., (1997) and Yoo et al., (1995) reported that RNA transcripts from cDNA clones of HCV-1 (genotype 1a) and HCV-N (genotype 1b), respectively, resulted in viral replication after transfection into human hepatoma cell lines. Unfortunately, the viability of these clones was not tested in vivo and concerns were raised about the infectivity of these cDNA clones in vitro (Fausto, 1997). In addition, both clones did not contain the terminal 98 conserved nucleotides at the very 3' end of the UTR.

Kolykhalov et al., (1997) and Yanagi et al. (1997) reported the derivation from HCV strain H77 (which is genotype 1a) of cDNA clones of HCV that are infectious for chimpanzees. However, while these infectious clones will aid in studying HCV replication and pathogenesis and will provide an important tool for development of in vitro replication and propagation systems, it is important to have infectious clones of more than one genotype given the extensive genetic heterogeneity of HCV and the potential

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impact of such heterogeneity on the development of effective therapies and vaccines for HCV.

# Summary Of The Invention

The present invention relates to nucleic acid sequences which comprise the genome of infectious hepatitis C viruses and in particular, nucleic acid sequences which comprise the genome of infectious hepatitis C viruses of genotype 1a and 1b strains. It is therefore an object of the invention to provide nucleic acid sequences which encode infectious hepatitis C viruses. Such nucleic acid sequences are referred to throughout the application as "infectious nucleic acid sequences".

For the purposes of this application, nucleic acid sequence refers to RNA, DNA, cDNA or any variant thereof capable of directing host organism synthesis of a hepatitis C virus polypeptide. It is understood that nucleic acid sequence encompasses nucleic acid sequences, which due to degeneracy, encode the same polypeptide sequence as the nucleic acid sequences described herein.

The invention also relates to the use of the infectious nucleic acid sequences to produce chimeric genomes consisting of portions of the open reading frames of infectious nucleic acid sequences of other genotypes (including, but not limited to, genotypes 1, 2, 3, 4, 5 and 6) and subtypes (including, but not limited to, subtypes 1a, 1b, 2a, 2b, 2c, 3a 4a-4f, 5a and 6a) of HCV. For example infectious nucleic acid sequence of the 1a and 1b strains H77 and HC-J4, respectively, described herein

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can be used to produce chimeras with sequences from the genomes of other strains of HCV from different genotypes or subtypes. Nucleic acid sequences which comprise sequence from the open-reading frames of 2 or more HCV genotypes or subtypes are designated "chimeric nucleic acid sequences".

The invention further relates to mutations of the infectious nucleic acid sequences of the invention where mutation includes, but is not limited to, point mutations, deletions and insertions. In one embodiment, a gene or fragment thereof can be deleted to determine the effect of the deleted gene or genes on the properties of the encoded virus such as its virulence and its ability to replicate. In an alternative embodiment, a mutation may be introduced into the infectious nucleic acid sequences to examine the effect of the mutation on the properties of the virus in the host cell.

The invention also relates to the introduction of mutations or deletions into the infectious nucleic acid sequences in order to produce an attenuated hepatitis C virus suitable for vaccine development.

The invention further relates to the use of the infectious nucleic acid sequences to produce attenuated viruses via passage in vitro or in vivo of the viruses produced by transfection of a host cell with the infectious nucleic acid sequence.

The present invention also relates to the use of the nucleic acid sequences of the invention or fragments thereof in the production of polypeptides where "nucleic acid sequences of the invention" refers to infectious

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nucleic acid sequences, mutations of infectious nucleic acid sequences, chimeric nucleic acid sequences and sequences which comprise the genome of attenuated viruses produced from the infectious nucleic acid sequences of the invention. The polypeptides of the invention, especially structural polypeptides, can serve as immunogens in the development of vaccines or as antigens in the development of diagnostic assays for detecting the presence of HCV in biological samples.

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The invention therefore also relates to vaccines for use in immunizing mammals especially humans against hepatitis C. In one embodiment, the vaccine comprises one or more polypeptides made from a nucleic acid sequence of the invention or fragment thereof. In a second embodiment, the vaccine comprises a hepatitis C virus produced by transfection of host cells with the nucleic acid sequences of the invention.

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The present invention therefore relates to methods for preventing hepatitis C in a mammal. In one embodiment the method comprises administering to a mammal a polypeptide or polypeptides encoded by a nucleic acid sequence of the invention in an amount effective to induce protective immunity to hepatitis C. In another embodiment, the method of prevention comprises administering to a mammal a hepatitis C virus of the invention in an amount effective to induce protective immunity against hepatitis C.

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In yet another embodiment, the method of protection comprises administering to a mammal a nucleic acid sequence of the invention or a fragment thereof in an

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amount effective to induce protective immunity against hepatitis C.

The invention also relates to hepatitis C viruses produced by host cells transfected with the nucleic acid sequences of the present invention.

The invention therefore also provides pharmaceutical compositions comprising the nucleic acid sequences of the invention and/or their encoded hepatitis C viruses. The invention further provides pharmaceutical compositions comprising polypeptides encoded by the nucleic acid sequences of the invention or fragments thereof. The pharmaceutical compositions of the invention may be used prophylactically or therapeutically.

The invention also relates to antibodies to the hepatitis C viruses of the invention or their encoded polypeptides and to pharmaceutical compositions comprising these antibodies.

The present invention further relates to polypeptides encoded by the nucleic acid sequences of the invention fragments thereof. In one embodiment, said polypeptide or polypeptides are fully or partially purified from hepatitis C virus produced by cells transfected with nucleic acid sequence of the invention. In another embodiment, the polypeptide or polypeptides are produced recombinantly from a fragment of the nucleic acid sequences of the invention. In yet another embodiment, the polypeptides are chemically synthesized.

The invention also relates to the use of the nucleic acid sequences of the invention to identify cell

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lines capable of supporting the replication of HCV <u>in</u> <u>vitro</u>.

The invention further relates to the use of the nucleic acid sequences of the invention or their encoded proteases (e.g. NS3 protease) to develop screening assays to identify antiviral agents for HCV.

# Brief Description Of Figures

Figure 1 shows a strategy for constructing fulllength cDNA clones of HCV strain H77. The long PCR products amplified with H1 and H9417R primers were cloned directly into pGEM-9zf(-) after digestion with Not I and Xba I (pH21, and pH50,). Next, the 3' UTR was cloned into both  $pH21_I$  and  $pH50_I$  after digestion with Afl II and Xba I (pH21 and pH50). pH21 was tested for infectivity in a chimpanzee. To improve the efficiency of cloning, we constructed a cassette vector with consensus 5' and 3' termini of H77. This cassette vector (pCV) was obtained by cutting out the BamHI fragment (nts 1358- 7530 of the H77 genome) from pH50, followed by religation. Finally, the long PCR products of H77 amplified with primers H1 and H9417R (H product) or primers A1 and H9417R (A product) were cloned into pCV after digestion with Age I and Afl II or with Pin AI and Bfr I. The latter procedure yielded multiple complete cDNA clones of strain H77 of HCV.

Figure 2 shows the results of gel electrophoresis of long RT-PCR amplicons of the entire ORF of H77 and the transcription mixture of the infectious clone of H77. The complete ORF was amplified by long RT-PCR with the primers H1 or A1 and H9417R from 10<sup>5</sup> GE of

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H77. A total of 10  $\mu$ g of the consensus chimeric clone (pCV-H77C) linearized with Xba I was transcribed in a 100  $\mu$ l reaction with T7 RNA polymerase. Five  $\mu$ l of the transcription mixture was analyzed by gel electrophoresis and the remainder of the mixture was injected into a chimpanzee. Lane 1, molecular weight marker; lane 2, products amplified with primers H1 and H9417R; lane 3,

products amplified with primers Al and H9417R; lane 4,

transcription mixture containing the RNA transcripts and linearized clone pCV-H77C (12.5 kb).

Figure 3 is a diagram of the genome organization of HCV strain H77 and the genetic heterogeneity of individual full-length clones compared with the consensus sequence of H77. Solid lines represent aa changes. Dashed lines represent silent mutations. A \* in pH21 represents a point mutation at nt 58 in the 5' UTR. the ORF, the consensus chimeric clone pCV-H77C had 11 nt differences [at positions 1625 ( $C \rightarrow T$ ), 2709 ( $T \rightarrow C$ ), 3380  $(A\rightarrow G)$ , 3710  $(C\rightarrow T)$ , 3914  $(G\rightarrow A)$ , 4463  $(T\rightarrow C)$ , 5058  $(C\rightarrow T)$ , 5834 (C $\rightarrow$ T), 6734 (T $\rightarrow$ C), 7154 (C $\rightarrow$ T), and 7202 (T $\rightarrow$ C)] and one as change (F  $\rightarrow$  L at as 790) compared with the consensus sequence of H77. This clone was infectious. Clone pH21 and pCV-H11 had 19 nts (7 aa) and 64 nts (21 aa) differences respectively, compared with the consensus sequence of H77. These two clones were not infectious. A single point mutation in the 3' UTR at nucleotide 9406 (G→A) introduced to create an Afl II cleavage site is not shown.

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Figures 4A-4F show the complete nucleotide sequence of a H77C clone produced according to the present invention and Figures 4G-4H show the amino acid sequence encoded by the H77C clone.

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Figure 5 shows an agarose gel of long RT-PCR amplicons and transcription mixtures. Lanes 1 and 4:

Molecular weight marker (Lambda/HindIII digest). Lanes 2 and 3: RT-PCR amplicons of the entire ORF of HC-J4. Lane 5: pCV-H77C transcription control (Yanagi et al., 1997).

Lanes 6, 7, and 8: 1/40 of each transcription mixture of pCV-J4L2S, pCV-J4L4S and pCV-J4L6S, respectively, which was injected into the chimpanzee.

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Figure 6 shows the strategy utilized for the construction of full-length cDNA clones of HCV strain HC-J4. The long PCR products were cloned as two separate fragments (L and S) into a cassette vector (pCV) with fixed 5' and 3' termini of HCV (Yanagi et al., 1997). Full-length cDNA clones of HC-J4 were obtained by inserting the L fragment from three pCV-J4L clones into three identical pCV-J4S9 clones after digestion with PinAI (isoschizomer of AgeI) and BfrI (isoschizomer of AfIII).

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Figure 7 shows amino acid positions with a quasispecies of HC-J4 in the acute phase plasma pool obtained from an experimentally infected chimpanzee.

Cons-p9: consensus amino acid sequence deduced from analysis of nine L fragments and nine S fragments (see Fig. 6). Cons-D: consensus sequence derived from direct sequencing of the PCR product. A, B, C: groups of similar viral species. Dot: amino acid identical to that in Cons-

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p9. Capital letter: amino acid different from that in Cons-p9. Cons-F: composite consensus amino acid sequence combining Cons-p9 and Cons-D. Boxed amino acid: different from that in Cons-F. Shaded amino acid: different from that in all species A sequences. An \*: defective ORF due to a nucleotide deletion (clone L1, aa 1097) or insertion (clone L7, aa 2770). Diagonal lines: fragments used to construct the infectious clone.

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Figure 8 shows comparisons (percent difference) of nucleotide (nts. 156 - 8935) and predicted amino acid sequences (aa 1 - 2864) of L clones (species A, B, and C, this study), HC-J4/91 (Okamoto et al., 1992b) and HC-J4/83 (Okamoto et al., 1992b). Differences among species A sequences and among species B sequences are shaded.

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Figure 9 shows UPGMA ("unweighted pair group method with arithmetic mean") trees of HC-J4/91 (Okamoto et al., 1992b), HC-J4/83 (Okamoto et al., 1992b), two prototype strains of genotype 1b (HCV-J, Kato et al., 1990; HCV-BK, Takamizawa et al., 1991), and L clones (this study).

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Figure 10 shows the alignment of the HVR1 and HVR2 amino acid sequences of the E2 sequences of nine L clones of HC-J4 (species A, B, and C) obtained from an early acute phase plasma pool of an experimentally infected chimpanzee compared with the sequences of eight clones (HC-J4/91-20 through HC-J4/91-27, Okamoto et al., 1992b) derived from the inoculum. Dot: an amino acid identical to that in the top line. Capital letters: amino acid different from that in the top line.

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Figure 11 shows the alignment of the 5' UTR and the 3' UTR sequences of infectious clones of genotype la (pCV-H77C) and 1b (pCV-J4L6S). Top line: consensus sequence of the indicated strain. Dot: identity with consensus sequence. Capital letter: different from the consensus sequence. Dash: deletion. Underlined: PinAI and BfrI cleavage site. Numbering corresponds to the HCV sequence of pCV-J4L6S.

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Figure 12 shows a comparison of individual full-length cDNA clones of the ORF of HCV strain HC-J4 with the consensus sequence (see Fig. 7). Solid lines: amino acid changes. Dashed lines: silent mutations. Clone pCV-J4L6S was infectious in vivo whereas clones pCV-J4L2S and pCV-J4L4S were not.

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Figure 13 shows biochemical (ALT levels) and PCR analyses of a chimpanzee following percutaneous intrahepatic transfection with RNA transcripts of the infectious clone of pCV-J4L2S, pCV-J4L4S and pCV-J4L6S. The ALT serum enzyme levels were measured in units per liter (u/l). For the PCR analysis, "HCV RNA" represented by an open rectangle indicates a serum sample that was negative for HCV after nested PCR; "HCV RNA" represented by a closed rectangle indicates that the serum sample was positive for HCV and HCV GE titer on the right-hand y-axis represents genome equivalents.

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Figures 14A-14F show the nucleotide sequence of the infectious clone of genotype 1b strain HC-J4 and Figures 14G-14H show the amino acid sequence encoded by the HC-J4 clone.

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Figure 15 shows the strategy for constructing a chimeric HCV clone designated pH77CV-J4 which contains the nonstructural region of the infectious clone of genotype la strain H77 and the structural region of the infectious clone of genotype lb strain HC-J4.

Figures 16A-16F show the nucleotide sequence of the chimeric 1a/1b clone pH77CV-J4 of Figure 15 and Figures 16G-16H show the amino acid sequence encoded by the chimeric 1a/1b clone.

Figures 17A and 17B show the sequence of the 3' untranslated region remaining in various 3' deletion mutants of the 1a infectious clone pCV-H77C and the strategy utilized in constructing each 3' deletion mutant (Figures 17C-17G).

Of the seven deletion mutants shown, two (pCV-H77C(-98X) and (-42X)) have been constructed and tested for infectivity in chimpanzees (see Figures 17A and 17C) and the other six are to be constructed and tested for infectivity as described in Figures 17D-17G.

Figures 18A and 18B show biochemical (ALT levels), PCR (HCV RNA and HCV GE titer), serological (anti-HCV) and histopathological (Fig. 18B only) analyses of chimpanzees 1494 (Fig. 18A) and 1530 (Fig. 18B) following transfection with the infectious cDNA clone pCV-H77C.

The ALT serum enzyme levels were measured in units per ml (u/l). For the PCR analysis, "HCV RNA" represented by an open rectangle indicates a serum sample that was negative for HCV after nested PCR; "HCV RNA" represented by a closed rectangle indicates that the serum

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sample was positive for HCV; and HCV GE titer on the right-hand y-axis represents genome equivalents.

The bar marked "anti-HCV" indicates samples that were positive for anti-HCV antibodies as determined by commercial assays. The histopathology scores in Figure 18B correspond to no histopathology (O), mild hepatitis (O) and moderate to severe hepatitis (O).

# DESCRIPTION OF THE INVENTION

The present invention relates to nucleic acid sequences which comprise the genome of an infectious hepatitis C virus. More specifically, the invention relates to nucleic acid sequences which encode infectious hepatitis C viruses of genotype 1a and 1b strains. In one embodiment, the infectious nucleic acid sequence of the invention has the sequence shown in Figures 4A-4F of this application. In another embodiment, the infectious nucleic acid sequence has the sequence shown in Figures 14A-14F and is contained in a plasmid construct deposited with the American Type Culture Collection (ATCC) on

The invention also relates to "chimeric nucleic acid sequences" where the chimeric nucleic acid sequences consist of open-reading frame sequences taken from infectious nucleic acid sequences of hepatitis C viruses of different genotypes or subtypes.

January 26, 1998 and having ATCC accession number 209596.

In one embodiment, the chimeric nucleic acid sequence consists of sequence from the genome of an HCV strain belonging to one genotype or subtype which encodes structural polypeptides and sequence of an HCV strain

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belonging to another genotype strain or subtype which encodes nonstructural polypeptides. Such chimeras can be produced by standard techniques of restriction digestion, PCR amplification and subcloning known to those of ordinary skill in the art.

In a preferred embodiment, the sequence encoding nonstructural polypeptides is from an infectious nucleic acid sequence encoding a genotype 1a strain where the construction of a chimeric 1a/1b nucleic acid sequence is described in Example 9 and the chimeric 1a/1b nucleic acid sequence is shown in Figures 16A-16F. It is believed that the construction of such chimeric nucleic acid sequences will be of importance in studying the growth and virulence properties of hepatitis C virus and in the production of hepatitis C viruses suitable to confer protection against multiple genotypes of HCV. For example, one might produce a "multivalent" vaccine by putting epitopes from several genotypes or subtypes into one clone. Alternatively one might replace just a single gene from an infectious sequence with the corresponding gene from the genomic sequence of a strain from another genotype or subtype or create a chimeric gene which contains portions of a gene from two genotypes or subtypes. Examples of genes which could be replaced or which could be made chimeric,

include, but are not limited to, the E1, E2 and NS4 genes.

The invention further relates to mutations of the infectious nucleic acid sequences where "mutations" includes, but is not limited to, point mutations, deletions and insertions. Of course, one of ordinary skill in the art would recognize that the size of the

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insertions would be limited by the ability of the resultant nucleic acid sequence to be properly packaged within the virion. Such mutation could be produced by techniques known to those of skill in the art such as site-directed mutagenesis, fusion PCR, and restriction digestion followed by religation.

In one embodiment, mutagenesis might be undertaken to determine sequences that are important for viral properties such as replication or virulence. example, one may introduce a mutation into the infectious nucleic acid sequence which eliminates the cleavage site between the NS4A and NS4B polypeptides to examine the effects on viral replication and processing of the polypeptide. Alternatively, one or more of the 3 amino acids encoded by the infectious 1b nucleic acid sequence shown in Figures 14A-14F which differ from the HC-J4 consensus sequence may be back mutated to the corresponding amino acid in the HC-J4 consensus sequence to determine the importance of these three amino acid changes to infectivity or virulence. In yet another embodiment, one or more of the amino acids from the noninfectious 1b clones pCV-J4L2S and pCV-J4L4S which differ from the consensus sequence may be introduced into the infectious 1b sequence shown in Figures 14A-14F.

In yet another example, one may delete all or part of a gene or of the 5' or 3' nontranslated region contained in an infectious nucleic acid sequence and then transfect a host cell (animal or cell culture) with the mutated sequence and measure viral replication in the host by methods known in the art such as RT-PCR. Preferred

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genes include, but are not limited to, the P7, NS4B and NS5A genes. Of course, those of ordinary skill in the art will understand that deletion of part of a gene, preferably the central portion of the gene, may be preferable to deletion of the entire gene in order to conserve the cleavage site boundaries which exist between proteins in the HCV polyprotein and which are necessary for proper processing of the polyprotein.

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In the alternative, if the transfection is into a host animal such as a chimpanzee, one can monitor the virulence phenotype of the virus produced by transfection of the mutated infectious nucleic acid sequence by methods known in the art such as measurement of liver enzyme levels (alanine aminotransferase (ALT) or isocitrate dehydrogenase (ICD)) or by histopathology of liver biopsies. Thus, mutations of the infectious nucleic acid sequences may be useful in the production of attenuated HCV strains suitable for vaccine use.

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The invention also relates to the use of the infectious nucleic acid sequences of the present invention to produce attenuated viral strains via passage <u>in vitro</u> or <u>in vivo</u> of the virus produced by transfection with the infectious nucleic acid sequences.

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The present invention therefore relates to the use of the nucleic acid sequences of the invention to identify cell lines capable of supporting the replication of HCV.

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In particular, it is contemplated that the mutations of the infectious nucleic acid sequences of the invention and the production of chimeric sequences as

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discussed above may be useful in identifying sequences critical for cell culture adaptation of HCV and hence, may be useful in identifying cell lines capable of supporting HCV replication.

Transfection of tissue culture cells with the nucleic acid sequences of the invention may be done by methods of transfection known in the art such as electroporation, precipitation with DEAE-Dextran or calcium phosphate or liposomes.

In one such embodiment, the method comprises the growing of animal cells, especially human cells, in vitro and transfecting the cells with the nucleic acid of the invention, then determining if the cells show indicia of HCV infection. Such indicia include the detection of viral antigens in the cell, for example, by immunofluorescent procedures well known in the art; the detection of viral polypeptides by Western blotting using antibodies specific therefor; and the detection of newly transcribed viral RNA within the cells via methods such as RT-PCR. The presence of live, infectious virus particles following such tests may also be shown by injection of cell culture medium or cell lysates into healthy, susceptible animals, with subsequent exhibition of the symptoms of HCV infection.

Suitable cells or cell lines for culturing HCV include, but are not limited to, lymphocyte and hepatocyte cell lines known in the art.

Alternatively, primary hepatocytes can be cultured, and then infected with HCV; or, the hepatocyte cultures could be derived from the livers of infected

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chimpanzees. In addition, various immortalization methods known to those of ordinary skill in the art can be used to obtain cell-lines derived from hepatocyte cultures. For example, primary hepatocyte cultures may be fused to a variety of cells to maintain stability.

The present invention further relates to the <u>in</u>

<u>vitro</u> and <u>in vivo</u> production of hepatitis C viruses from

the nucleic acid sequences of the invention.

In one embodiment, the sequences of the invention can be inserted into an expression vector that functions in eukaryotic cells. Eukaryotic expression vectors suitable for producing high efficiency gene transfer in vivo are well known to those of ordinary skill in the art and include, but are not limited to, plasmids, vaccinia viruses, retroviruses, adenoviruses and adenoassociated viruses.

In another embodiment, the sequences contained in the recombinant expression vector can be transcribed <u>in vitro</u> by methods known to those of ordinary skill in the art in order to produce RNA transcripts which encode the hepatitis C viruses of the invention. The hepatitis C viruses of the invention may then be produced by transfecting cells by methods known to those of ordinary skill in the art with either the <u>in vitro</u> transcription mixture containing the RNA transcripts (see Example 4) or with the recombinant expression vectors containing the nucleic acid sequences described herein.

The present invention also relates to the construction of cassette vectors useful in the cloning of viral genomes wherein said vectors comprise a nucleic acid

sequence to be cloned, and said vector reading in the correct phase for the expression of the viral nucleic acid to be cloned. Such a cassette vector will, of course, also possess a promoter sequence, advantageously placed 5 upstream of the sequence to be expressed. Cassette vectors according to the present invention are constructed according to the procedure described in Figure 1, for example, starting with plasmid pCV. Of course, the DNA to be inserted into said cassette vector can be derived from 10 any virus, advantageously from HCV, and most advantageously from the H77 strain of HCV. The nucleic acid to be inserted according to the present invention can, for example, contain one or more open reading frames 15 of the virus, for example, HCV. The cassette vectors of the present invention may also contain, optionally, one or more expressible marker genes for expression as an indication of successful transfection and expression of the nucleic acid sequences of the vector. To insure 20 expression, the cassette vectors of the present invention will contain a promoter sequence for binding of the appropriate cellular RNA polymerase, which will depend on the cell into which the vector has been introduced. 25 example, if the host cell is a bacterial cell, then said promoter will be a bacterial promoter sequence to which the bacterial RNA polymerases will bind.

The hepatitis C viruses produced from the sequences of the invention may be purified or partially purified from the transfected cells by methods known to those of ordinary skill in the art. In a preferred embodiment, the viruses are partially purified prior to

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their use as immunogens in the pharmaceutical compositions and vaccines of the present invention.

The present invention therefore relates to the use of the hepatitis C viruses produced from the nucleic acid sequences of the invention as immunogens in live or killed (e.g., formalin inactivated) vaccines to prevent hepatitis C in a mammal.

In an alternative embodiment, the immunogen of the present invention may be an infectious nucleic acid sequence, a chimeric nucleic acid sequence, or a mutated infectious nucleic acid sequence which encodes a hepatitis C virus. Where the sequence is a cDNA sequence, the cDNAs and their RNA transcripts may be used to transfect a mammal by direct injection into the liver tissue of the mammal as described in the Examples.

Alternatively, direct gene transfer may be accomplished via administration of a eukaryotic expression vector containing a nucleic acid sequence of the invention.

In yet another embodiment, the immunogen may be a polypeptide encoded by the nucleic acid sequences of the invention. The present invention therefore also relates to polypeptides produced from the nucleic acid sequences of the invention or fragments thereof. In one embodiment, polypeptides of the present invention can be recombinantly produced by synthesis from the nucleic acid sequences of the invention or isolated fragments thereof, and purified, or partially purified, from transfected cells using methods already known in the art. In an alternative embodiment, the polypeptides may be purified or partially

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purified from viral particles produced via transfection of a host cell with the nucleic acid sequences of the invention. Such polypeptides might, for example, include either capsid or envelope polypeptides prepared from the sequences of the present invention.

When used as immunogens, the nucleic acid sequences of the invention, or the polypeptides or viruses produced therefrom, are preferably partially purified prior to use as immunogens in pharmaceutical compositions and vaccines of the present invention. When used as a vaccine, the sequences and the polypeptide and virus products thereof, can be administered alone or in a suitable diluent, including, but not limited to, water, saline, or some type of buffered medium. The vaccine according to the present invention may be administered to an animal, especially a mammal, and most especially a human, by a variety of routes, including, but not limited to, intradermally, intramuscularly, subcutaneously, or in any combination thereof.

Suitable amounts of material to administer for prophylactic and therapeutic purposes will vary depending on the route selected and the immunogen (nucleic acid, virus, polypeptide) administered. One skilled in the art will appreciate that the amounts to be administered for any particular treatment protocol can be readily determined without undue experimentation. The vaccines of the present invention may be administered once or periodically until a suitable titer of anti-HCV antibodies appear in the blood. For an immunogen consisting of a nucleic acid sequence, a suitable amount of nucleic acid

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sequence to be used for prophylactic purposes might be expected to fall in the range of from about 100  $\mu g$  to about 5 mg and most preferably in the range of from about 500  $\mu g$  to about 2mg. For a polypeptide, a suitable amount to use for prophylactic purposes is preferably 100 ng to 100  $\mu g$  and for a virus  $10^2$  to  $10^6$  infectious doses. Such administration will, of course, occur prior to any sign of HCV infection.

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A vaccine of the present invention may be employed in such forms as capsules, liquid solutions, suspensions or elixirs for oral administration, or sterile liquid forms such as solutions or suspensions. Any inert carrier is preferably used, such as saline or phosphatebuffered saline, or any such carrier in which the HCV of the present invention can be suitably suspended. vaccines may be in the form of single dose preparations or in multi-dose flasks which can be utilized for massvaccination programs of both animals and humans. purposes of using the vaccines of the present invention reference is made to Remington's Pharmaceutical Sciences, Mack Publishing Co., Easton, Pa., Osol (Ed.) (1980); and New Trends and Developments in Vaccines, Voller et al. (Eds.), University Park Press, Baltimore, Md. (1978), both of which provide much useful information for preparing and using vaccines. Of course, the polypeptides of the present invention, when used as vaccines, can include, as part of the composition or emulsion, a suitable adjuvant, such as alum (or aluminum hydroxide) when humans are to be vaccinated, to further stimulate production of antibodies by immune cells. When nucleic acids or viruses are used

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for vaccination purposes, other specific adjuvants such as CpG motifs (Krieg, A.K. et al.(1995) and (1996)), may prove useful.

When the nucleic acids, viruses and polypeptides of the present invention are used as vaccines or inocula, they will normally exist as physically discrete units suitable as a unitary dosage for animals, especially mammals, and most especially humans, wherein each unit will contain a predetermined quantity of active material calculated to produce the desired immunogenic effect in association with the required diluent. The dose of said vaccine or inoculum according to the present invention is administered at least once. In order to increase the antibody level, a second or booster dose may be administered at some time after the initial dose. need for, and timing of, such booster dose will, of course, be determined within the sound judgment of the administrator of such vaccine or inoculum and according to sound principles well known in the art. For example, such booster dose could reasonably be expected to be advantageous at some time between about 2 weeks to about 6 months following the initial vaccination. Subsequent doses may be administered as indicated.

The nucleic acid sequences, viruses and polypeptides of the present invention can also be administered for purposes of therapy, where a mammal, especially a primate, and most especially a human, is already infected, as shown by well known diagnostic measures. When the nucleic acid sequences, viruses or polypeptides of the present invention are used for such

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therapeutic purposes, much of the same criteria will apply as when it is used as a vaccine, except that inoculation will occur post-infection. Thus, when the nucleic acid sequences, viruses or polypeptides of the present invention are used as therapeutic agents in the treatment of infection, the therapeutic agent comprises a pharmaceutical composition containing a sufficient amount of said nucleic acid sequences, viruses or polypeptides so as to elicit a therapeutically effective response in the organism to be treated. Of course, the amount of pharmaceutical composition to be administered will, as for vaccines, vary depending on the immunogen contained therein (nucleic acid, polypeptide, virus) and on the route of administration.

The therapeutic agent according to the present invention can thus be administered by, subcutaneous, intramuscular or intradermal routes. One skilled in the art will certainly appreciate that the amounts to be administered for any particular treatment protocol can be readily determined without undue experimentation. Of course, the actual amounts will vary depending on the route of administration as well as the sex, age, and clinical status of the subject which, in the case of human patients, is to be determined with the sound judgment of the clinician.

The therapeutic agent of the present invention

can be employed in such forms as capsules, liquid solutions, suspensions or elixirs, or sterile liquid forms such as solutions or suspensions. Any inert carrier is preferably used, such as saline, phosphate-buffered

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saline, or any such carrier in which the HCV of the present invention can be suitably suspended. The therapeutic agents may be in the form of single dose preparations or in the multi-dose flasks which can be utilized for mass-treatment programs of both animals and humans. Of course, when the nucleic acid sequences, viruses or polypeptides of the present invention are used as therapeutic agents they may be administered as a single dose or as a series of doses, depending on the situation as determined by the person conducting the treatment.

The nucleic acids, polypeptides and viruses of the present invention can also be utilized in the production of antibodies against HCV. The term "antibody" is herein used to refer to immunoglobulin molecules and immunologically active portions of immunoglobulin molecules. Examples of antibody molecules are intact immunoglobulin molecules, substantially intact immunoglobulin molecules and portions of an immunoglobulin molecule, including those portions known in the art as Fab, F(ab')<sub>2</sub> and F(v) as well as chimeric antibody molecules.

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Thus, the polypeptides, viruses and nucleic acid sequences of the present invention can be used in the generation of antibodies that immunoreact (i.e., specific binding between an antigenic determinant-containing molecule and a molecule containing an antibody combining site such as a whole antibody molecule or an active portion thereof) with antigenic determinants on the surface of hepatitis C virus particles.

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The present invention therefore also relates to antibodies produced following immunization with the nucleic acid sequences, viruses or polypeptides of the present invention. These antibodies are typically produced by immunizing a mammal with an immunogen or vaccine to induce antibody molecules having immunospecificity for polypeptides or viruses produced in response to infection with the nucleic acid sequences of the present invention. When used in generating such antibodies, the nucleic acid sequences, viruses, or polypeptides of the present invention may be linked to some type of carrier molecule. The resulting antibody molecules are then collected from said mammal. Antibodies produced according to the present invention have the unique advantage of being generated in response to authentic, functional polypeptides produced according to the actual cloned HCV genome.

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The antibody molecules of the present invention may be polyclonal or monoclonal. Monoclonal antibodies are readily produced by methods well known in the art. Portions of immunoglobin molecules, such as Fabs, as well as chimeric antibodies, may also be produced by methods well known to those of ordinary skill in the art of generating such antibodies.

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The antibodies according to the present invention may also be contained in blood plasma, serum, hybridoma supernatants, and the like. Alternatively, the antibody of the present invention is isolated to the extent desired by well known techniques such as, for example, using DEAE Sephadex. The antibodies produced

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according to the present invention may be further purified so as to obtain specific classes or subclasses of antibody such as IgM, IgG, IgA, and the like. Antibodies of the IgG class are preferred for purposes of passive protection.

The antibodies of the present invention are useful in the prevention and treatment of diseases caused by hepatitis C virus in animals, especially mammals, and most especially humans.

In providing the antibodies of the present invention to a recipient mammal, preferably a human, the dosage of administered antibodies will vary depending on such factors as the mammal's age, weight, height, sex, general medical condition, previous medical history, and the like.

In general, it will be advantageous to provide the recipient mammal with a dosage of antibodies in the range of from about 1 mg/kg body weight to about 10 mg/kg body weight of the mammal, although a lower or higher dose may be administered if found desirable. Such antibodies will normally be administered by intravenous or intramuscular route as an inoculum. The antibodies of the present invention are intended to be provided to the recipient subject in an amount sufficient to prevent, lessen or attenuate the severity, extent or duration of any existing infection.

The antibodies prepared by use of the nucleic acid sequences, viruses or polypeptides of the present invention are also highly useful for diagnostic purposes. For example, the antibodies can be used as in vitro

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diagnostic agents to test for the presence of HCV in biological samples taken from animals, especially humans. Such assays include, but are not limited to, radioimmunoassays, EIA, fluorescence, Western blot analysis and ELISAs. In one such embodiment, the biological sample is contacted with antibodies of the present invention and a labeled second antibody is used to detect the presence of HCV to which the antibodies are bound.

Such assays may be, for example, a direct protocol (where the labeled first antibody is immunoreactive with the antigen, such as, for example, a polypeptide on the surface of the virus), an indirect protocol (where a labeled second antibody is reactive with the first antibody), a competitive protocol (such as would involve the addition of a labeled antigen), or a sandwich protocol (where both labeled and unlabeled antibody are used), as well as other protocols well known and described in the art.

In one embodiment, an immunoassay method would utilize an antibody specific for HCV envelope determinants and would further comprise the steps of contacting a biological sample with the HCV-specific antibody and then detecting the presence of HCV material in the test sample using one of the types of assay protocols as described above. Polypeptides and antibodies produced according to the present invention may also be supplied in the form of a kit, either present in vials as purified material, or present in compositions and suspended in suitable diluents as previously described.

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In a preferred embodiment, such a diagnostic test kit for detection of HCV antigens in a test sample comprises in combination a series of containers, each container a reagent needed for such assay. Thus, one such container would contain a specific amount of HCV-specific antibody as already described, a second container would contain a diluent for suspension of the sample to be tested, a third container would contain a positive control and an additional container would contain a negative control. An additional container could contain a blank.

For all prophylactic, therapeutic and diagnostic uses, the antibodies of the invention and other reagents, plus appropriate devices and accessories, may be provided in the form of a kit so as to facilitate ready availability and ease of use.

The present invention also relates to the use of nucleic acid sequences and polypeptides of the present invention to screen potential antiviral agents for antiviral activity against HCV. Such screening methods are known by those of skill in the art. Generally, the antiviral agents are tested at a variety of concentrations, for their effect on preventing viral replication in cell culture systems which support viral replication, and then for an inhibition of infectivity or of viral pathogenicity (and a low level of toxicity) in an animal model system.

In one embodiment, animal cells (especially human cells) transfected with the nucleic acid sequences of the invention are cultured <u>in vitro</u> and the cells are treated with a candidate antiviral agent (a chemical,

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peptide etc.) for antiviral activity by adding the candidate agent to the medium. The treated cells are then exposed, possibly under transfecting or fusing conditions known in the art, to the nucleic acid sequences of the present invention. A sufficient period of time would then be allowed to pass for infection to occur, following which the presence or absence of viral replication would be determined versus untreated control cells by methods known to those of ordinary skill in the art. Such methods include, but are not limited to, the detection of viral antigens in the cell, for example, by immunofluorescent procedures well known in the art; the detection of viral polypeptides by Western blotting using antibodies specific therefor; the detection of newly transcribed viral RNA within the cells by RT-PCR; and the detection of the presence of live, infectious virus particles by injection of cell culture medium or cell lysates into healthy, susceptible animals, with subsequent exhibition of the symptoms of HCV infection. A comparison of results obtained for control cells (treated only with nucleic acid sequence) with those obtained for treated cells (nucleic acid sequence and antiviral agent) would indicate, the degree, if any, of antiviral activity of the candidate antiviral agent. Of course, one of ordinary skill in the art would readily understand that such cells can be treated with the candidate antiviral agent either before or after exposure to the nucleic acid sequence of the present invention so as to determine what stage, or stages, of viral infection and replication said agent is effective against.

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In an alternative embodiment, a protease such as NS3 protease produced from a nucleic acid sequence of the invention may be used to screen for protease inhibitors which may act as antiviral agents. The structural and nonstructural regions of the HCV genome, including nucleotide and amino acid locations, have been determined, for example, as depicted in Houghton, M. (1996), Fig. 1; and Major, M.E. et al. (1997), Table 1.

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Such above-mentioned protease inhibitors may take the form of chemical compounds or peptides which mimic the known cleavage sites of the protease and may be screened using methods known to those of skill in the art (Houghton, M. (1996) and Major, M.E. et al. (1997)). For example, a substrate may be employed which mimics the protease's natural substrate, but which provides a detectable signal (e.g. by fluorimetric or colorimetric methods) when cleaved. This substrate is then incubated with the protease and the candidate protease inhibitor under conditions of suitable pH, temperature etc. to detect protease activity. The proteolytic activities of the protease in the presence or absence of the candidate inhibitor are then determined.

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In yet another embodiment, a candidate antiviral agent (such as a protease inhibitor) may be directly assayed <u>in vivo</u> for antiviral activity by administering the candidate antiviral agent to a chimpanzee transfected with a nucleic acid sequence of the invention and then measuring viral replication <u>in vivo</u> via methods such as RT-PCR. Of course, the chimpanzee may be treated with the candidate agent either before or after transfection with

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the infectious nucleic acid sequence so as to determine what stage, or stages, of viral infection and replication the agent is effective against.

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The invention also provides that the nucleic acid sequences, viruses and polypeptides of the invention may be supplied in the form of a kit, alone or in the form of a pharmaceutical composition.

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All scientific publication and/or patents cited herein are specifically incorporated by reference. The following examples illustrate various aspects of the invention but are in no way intended to limit the scope thereof.

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### **EXAMPLES**

## MATERIALS AND METHODS For Examples 1-4

Collection of Virus

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Hepatitis C virus was collected and used as a source for the RNA used in generating the cDNA clones according to the present invention. Plasma containing strain H77 of HCV was obtained from a patient in the acute phase of transfusion-associated non-A, non-B hepatitis (Feinstone et al (1981)). Strain H77 belongs to genotype la of HCV (Ogata et al (1991), Inchauspe et al (1991)). The consensus sequence for most of its genome has been determined (Kolyakov et al (1996), Ogata et al (1991), Inchauspe et al (1991) and Farci et al (1996)).

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## RNA Purification

Viral RNA was collected and purified by conventional means. In general, total RNA from 10  $\mu$ l of H77 plasma was extracted with the TRIzol system (GIBCO BRL). The RNA pellet was resuspended in 100  $\mu$ l of 10 mM dithiothreitol (DTT) with 5% (vol/vol) RNasin (20 - 40 units/ $\mu$ l) (available from Promega) and 10  $\mu$ l aliquots were stored at -80°C. In subsequent experiments RT-PCR was performed on RNA equivalent to 1  $\mu$ l of H77 plasma, which contained an estimated 10<sup>5</sup> genome equivalents (GE) of HCV (Yanaqi et al (1996)).

Primers used in the RT-PCR process were deduced from the genomic sequences of strain H77 according to procedures already known in the art (see above) or else were determined specifically for use herein. The primers generated for this purpose are listed in Table 1.

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Table 1. Oligonucleotides used for PCR amplification of strain H77 of HCV

		Designation Sequence (5' → 3')*
5	H9261F	GGCTACAGCGGGGGAGACATTTATCACAGC
10	H3'X58R	TCATGCGGCTCACGGACCTTTCACAGCTAG
	H9282F	GTCCAAGCTTATCACAGCGTGTCTCATGCCCGGCCCCG
	H3'X45R	CGTCTCTAGAGGACCTTTCACAGCTAGCCGTGACTAGGG
	H9375F	TGAAGGTTGGGGTAAACACTCCGGCCTCTTAGGCCATT
	H3'X-35R	ACATGATCTGCAGAGAGGCCAGTATCAGCACTCTC
	H9386F	GTCCAAGCTTACGCGTAAACACTCCGGCCTCCTTAAGCCATTCCTG
	H3'X-38R	CGTCTCTAGACATGATCTGCAGAGAGGCCAGTATCAGCACTCTCTGC
	H1	TTTTTTTGCGGCCGCTAATACGACTCACTATAGCCAGCCCCCTGAT-
		GGGGGCGACACTCCACCATG
	A1	ACTGTCTTCACGCAGAAAGCGTCTAGCCAT
	H9417R	CGTCTCTAGACAGGAAATGGCTTAAGAGGCCGGAGTGTTTACC
	* HCV sequences are shown in plain text, non-HCV-specific	
15	sequences are shown in boldface and artificial cleavage sites	
	used for cDNA cloning are underlined. The core sequenceof the	
	T7 promoter in primer H1 is shown in italics.	
	Primers for long RT-PCR were size-purified.	

### cDNA Synthesis

The RNA was denatured at 65°C for 2 min, and cDNA synthesis was performed in a 20  $\mu$ l reaction volume with Superscript II reverse transcriptase (from GIBCO/BRL) at 42 °C for 1 hour using specific antisense primers as described previously (Tellier et al (1996)). The cDNA mixture was treated with RNase H and RNase T1 (GIBCO/BRL) for 20 min at 37 °C.

## Amplification and Cloning of the 3' UTR

The 3' UTR of strain H77 was amplified by PCR in two different assays. In both of these nested PCR reactions the first round of PCR was performed in a total volume of 50  $\mu$ l in 1 x buffer, 250  $\mu$ mol of each deoxynucleoside triphosphate (dNTP; Pharmacia), 20 pmol

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each of external sense and antisense primers, 1  $\mu$ l of the Advantage KlenTaq polymerase mix (from Clontech) and 2  $\mu$ l of the final cDNA reaction mixture. In the second round of PCR, 5  $\mu$ l of the first round PCR mixture was added to 5 45  $\mu$ l of PCR mixture prepared as already described. round of PCR (35 cycles), which was performed in a Perkin Elmer DNA thermal cycler 480, consisted of denaturation at 94 °C for 1 min (in 1st cycle 1 min 30 sec), annealing at 60°C for 1 min and elongation at 68°C for 2 min. 10 experiment a region from NS5B to the conserved region of the 3' UTR was amplified with the external primers H9261F and H3'X58R, and the internal primers H9282F and H3'X45R (Table 1). In another experiment, a segment of the 15 variable region to the very end of the 3' UTR was amplified with the external primers H9375F and H3'X-35R, and the internal primers H9386F and H3'X-38R (Table 1, Fig. 1). Amplified products were purified with QIAquick 20 PCR purification kit (from QIAGEN), digested with Hind III and Xba I (from Promega), purified by either gel electrophoresis or phenol/chloroform extraction, and then cloned into the multiple cloning site of plasmid pGEM-9zf(-) (Promega) or pUC19 (Pharmacia). Cloning of cDNA 25 into the vector was performed with T4 DNA ligase (Promega) by standard procedures.

# Amplification of Near Full-Length H77 Genomes by Long PCR

The reactions were performed in a total volume of 50  $\mu$ l in 1 x buffer, 250  $\mu$ mol of each dNTP, 10 pmol each of sense and antisense primers, 1  $\mu$ l of the Advantage

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KlenTaq polymerase mix and 2  $\mu$ l of the cDNA reaction mixture (Tellier et al (1996)). A single PCR round of 35 cycles was performed in a Robocycler thermal cycler (from Stratagene), and consisted of denaturation at 99 °C for 35 sec, annealing at 67 °C for 30 sec and elongation at 68 °C for 10 min during the first 5 cycles, 11 min during the next 10 cycles, 12 min during the following 10 cycles and 13 min during the last 10 cycles. To amplify the complete ORF of HCV by long RT-PCR we used the sense primers H1 or A1 deduced from the 5' UTR and the antisense primer H9417R deduced from the variable region of the 3' UTR (Table 1, Fig. 1).

The long PCR products amplified with H1 and

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### Construction of Full-Length H77 cDNA Clones

H9417R primers were cloned directly into pGEM-9zf(-) after digestion with Not I and Xba I (from Promega) (as per 20 Fig. 1). Two clones were obtained with inserts of the expected size, pH21, and pH50,. Next, the chosen 3' UTR was cloned into both pH21, and pH50, after digestion with Afl II and Xba I (New England Biolabs). DH5α competent 25 cells (GIBCO/BRL) were transformed and selected with LB agar plates containing 100  $\mu$ g/ml ampicillin (from SIGMA). Then the selected colonies were cultured in LB liquid containing ampicillin at 30°C for ~18-20 hrs 30 (transformants containing full-length or near full-length cDNA of H77 produced a very low yield of plasmid when cultured at 37 °C or for more than 24 hrs). After small scale preparation (Wizard Plus Minipreps DNA Purification

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Systems, Promega) each plasmid was retransformed to select a single clone, and large scale preparation of plasmid DNA was performed with a QIAGEN plasmid Maxi kit.

# 5 Cloning of Long RT-PCR Products Into a Cassette Vector

with consensus 5' and 3' termini of HCV strain H77 was constructed (Fig. 1). This cassette vector (pCV) was obtained by cutting out the BamHI fragment (nts 1358 - 7530 of the H77 genome) from pH50, followed by religation. Next, the long PCR products of H77 amplified with H1 and H9417R or A1 and H9417R primers were purified (Geneclean spin kit; BIO 101) and cloned into pCV after digestion with Age I and Afl II (New England Biolabs) or with Pin AI (isoschizomer of Age I) and Bfr I (isoschizomer of Afl II) (Boehringer Mannheim). Large scale preparations of the plasmids containing full-length cDNA of H77 were performed as described above.

# Construction of H77 Consensus Chimeric cDNA Clone

25 encoding the consensus amino acid sequence was constructed by making a chimera from four of the cDNA clones obtained above. This consensus chimera, pCV-H77C, was constructed in two ligation steps by using standard molecular procedures and convenient cleavage sites and involved first a two piece ligation and then a three piece ligation. Large scale preparation of pCV-H77C was performed as already described.

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### In Vitro Transcription

Plasmids containing the full-length HCV cDNA were linearized with Xba I (from Promega), and purified by phenol/chloroform extraction and ethanol precipitation. A 100  $\mu$ l reaction mixture containing 10  $\mu$ g of linearized plasmid DNA, 1 x transcription buffer, 1 mM ATP, CTP, GTP and UTP, 10mM DTT, 4% (v/v) RNasin (20-40 units/ $\mu$ l) and 2  $\mu$ l of T7 RNA polymerase (Promega) was incubated at 37 °C for 2 hrs. Five  $\mu$ l of the reaction mixture was analyzed by agarose gel electrophoresis followed by ethidium bromide staining. The transcription reaction mixture was diluted with 400  $\mu$ l of ice-cold phosphate-buffered saline without calcium or magnesium, immediately frozen on dry ice and stored at -80 °C. The final nucleic acid mixture was injected into chimpanzees within 24 hrs.

## Intrahepatic Transfection of Chimpanzees

Laparotomy was performed and aliquots from two transcription reactions were injected into 6 sites of the exposed liver (Emerson et al (1992). Serum samples were collected weekly from chimpanzees and monitored for liver enzyme levels and anti-HCV antibodies. Weekly samples of 100 µl of serum were tested for HCV RNA in a highly sensitive nested RT-PCR assay with AmpliTaq Gold (Perkin Elmer) (Yanagi et al (1996); Bukh et al (1992)). The genome titer of HCV was estimated by testing 10-fold serial dilutions of the extracted RNA in the RT-PCR assay (Yanagi et al (1996)). The two chimpanzees used in this

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study were maintained under conditions that met all requirements for their use in an approved facility.

The consensus sequence of the complete ORF from HCV genomes recovered at week 2 post inoculation (p.i) was determined by direct sequencing of PCR products obtained in long RT-PCR with primers A1 and H9417R followed by nested PCR of 10 overlapping fragments. The consensus sequence of the variable region of the 3' UTR was determined by direct sequencing of an amplicon obtained in nested RT-PCR as described above. Finally, we amplified selected regions independently by nested RT-PCR with AmpliTaq Gold.

### 15 <u>Sequence Analysis</u>

Both strands of DNA from PCR products, as well as plasmids, were sequenced with the ABI PRISM Dye Termination Cycle Sequencing Ready Reaction Kit using Taq DNA polymerase (Perkin Elmer) and about 100 specific sense and antisense sequence primers.

The consensus sequence of HCV strain H77 was determined in two different ways. In one approach, overlapping PCR products were directly sequenced, and amplified in nested RT-PCR from the H77 plasma sample. The sequence analyzed (nucleotides (nts) 35-9417) included the entire genome except the very 5' and 3' termini. In the second approach, the consensus sequence of nts 157-9384 was deduced from the sequences of 18 full-length cDNA clones.

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#### EXAMPLE 1

# Variability in the sequence of the 3' UTR of HCV strain H77

5 The heterogeneity of the 3' UTR was analyzed by cloning and sequencing of DNA amplicons obtained in nested 19 clones containing sequences of the entire RT-PCR. variable region, the poly U-UC region and the adjacent 19 nt of the conserved region, and 65 clones containing 10 sequences of the entire poly U-UC region and the first 63 nts of the conserved region were analyzed. This analysis confirmed that the variable region consisted of 43 nts, including two conserved termination codons (Han et al 15 (1992)). The sequence of the variable region was highly conserved within H77 since only 3 point mutations were found among the 19 clones analyzed. A poly U-UC region was present in all 84 clones analyzed. However, its length varied from 71-141 nts. The length of the poly U 20 region was 9-103 nts, and that of the poly UC region was 35-85 nts. The number of C residues increased towards the 3' end of the poly UC region but the sequence of this region is not conserved. The first 63 nts of the 25 conserved region were highly conserved among the clones analyzed, with a total of only 14 point mutations. confirm the validity of the analysis, the 3' UTR was reamplified directly from a full-length cDNA clone of HCV 30 (see below) by the nested-PCR procedure with the primers in the variable region and at the very 3' end of the HCV genome and cloned the PCR product. Eight clones had 1-7 nt deletions in the poly U region. Furthermore, although

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the C residues of the poly UC region were maintained, the spacing of these varied because of 1-2 nt deletions of U residues. These deletions must be artifacts introduced by PCR and such mistakes may have contributed to the heterogeneity originally observed in this region.

However, the conserved region of the 3' UTR was amplified correctly, suggesting that the deletions were due to difficulties in transcribing a highly repetitive sequence.

One of the 3' UTR clones was selected for engineering of full-length cDNA clones of H77. This clone had the consensus variable sequence except for a single point mutation introduced to create an Afl II cleavage site, a poly U-UC stretch of 81 nts with the most commonly observed UC pattern and the consensus sequence of the complete conserved region of 101 nts, including the distal 38 nts which originated from the antisense primer used in the amplification. After linearization with Xba I, the DNA template of this clone had the authentic 3' end.

#### EXAMPLE 2

# The Entire Open Reading Frame of H77 Amplified in One Round of Long RT-PCR

It had been previously demonstrated that a 9.25 kb fragment of the HCV genome from the 5' UTR to the 3' end of NS5B could be amplified from 10<sup>4</sup> GE (genome equivalents) of H77 by a single round of long RT-PCR (Tellier et al (1996a)). In the current study, by optimizing primers and cycling conditions, the entire ORF of H77 was amplified in a single round of long RT-PCR with primers from the 5' UTR and the variable region of the 3'

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UTR. In fact, 9.4 kb of the H77 genome (H product: from the very 5' end to the variable region of the 3' UTR) could be amplified from 10<sup>5</sup> GE or 9.3 kb (A product: from within the 5' UTR to the variable region of the 3' UTR) from 10<sup>4</sup> GE or 10<sup>5</sup> GE, in a single round of long RT-PCR (Fig. 2). The PCR products amplified from 10<sup>5</sup> GE of H77 were used for engineering full-length cDNA clones (see below).

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## EXAMPLE 3

Construction of Multiple Full-Length

cDNA Clones of H77 in a Single Step by

Cloning of Long RT-PCR Amplicons Directly

into a Cassette Vector with Fixed 5' and 3' Termini

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Direct cloning of the long PCR products (H), which contained a 5' T7 promoter, the authentic 5' end, the entire ORF of H77 and a short region of the 3' UTR, into pGEM-9zf(-) vector by Not I and Xba I digestion was first attempted. However, among the 70 clones examined all but two had inserts that were shorter than predicted. Sequence analysis identified a second Not I site in the majority of clones, which resulted in deletion of the nts past position 9221. Only two clones (pH21, and pH50,) were missing the second Not I site and had the expected 5' and 3' sequences of the PCR product. Therefore, full-length cDNA clones (pH21 and pH50) were constructed by inserting the chosen 3' UTR into pH21, and pH50, respectively. Sequence analysis revealed that clone pH21 had a complete full-length sequence of H77; this clone was tested for infectivity. The second clone, pH50, had one nt deletion

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in the ORF at position 6365; this clone was used to make a cassette vector.

The complete ORF was amplified by constructing a cassette vector with fixed 5' and 3' termini as an intermediate of the full-length cDNA clones. This vector (pCV) was constructed by digestion of clone pH50 with BamHI, followed by religation, to give a shortened plasmid readily distinguished from plasmids containing the fulllength insert. Attempts to clone long RT-PCR products (H) into pCV by Age I and Afl II yielded only 1 of 23 clones with an insert of the expected size. In order to increase the efficiency of cloning, we repeated the procedure but used Pin A I and Bfr I instead of the respective isoschizomers Age I and Afl II. By this protocol, 24 of 31 H clones and 30 of 35 A clones had the full-length cDNA of H77 as evaluated by restriction enzyme digestion. A total of 16 clones, selected at random, were each retransformed, and individual plasmids were purified and completely sequenced.

### EXAMPLE 4

Demonstration of Infectious Nature of Transcripts of a cDNA Clone Representing the Consensus Sequence of Strain H77

A consensus chimera was constructed from 4 of
the full-length cDNA clones with just 2 ligation steps.
The final construct, pCV-H77C, had 11 nt differences from
the consensus sequence of H77 in the ORF (Fig. 3).
However, 10 of these nucleotide differences represented
silent mutations. The chimeric clone differed from the

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consensus sequence at only one amino acid [L instead of F at position 790]. Among the 18 ORFs analyzed above, the F residue was found in 11 clones and the L residue in 7 clones. However, the L residue was dominant in other isolates of genotype 1a, including a first passage of H77 in a chimpanzee (Inchauspe et al (1991)).

chimeric clone of H77 intrahepatic transfection of a chimpanzee was performed. The pCV-H77C clone was linearized with Xba I and transcribed in vitro by T7 RNA polymerase (Fig. 2). The transcription mixture was next injected into 6 sites of the liver of chimpanzee 1530. The chimpanzee became infected with HCV as measured by detection of 10<sup>2</sup> GE/ml of viral genome at week 1 p.i. Furthermore, the HCV titer increased to 10<sup>4</sup> GE/ml at week 2 p.i., and reached 10<sup>6</sup> GE/ml by week 8 p.i. The viremic pattern observed in the early phase of the infection with the recombinant virus was similar to that observed in chimpanzees inoculated intravenously with strain H77 or other strains of HCV (Shimizu (1990)).

The sequence of the HCV genomes from the serum sample collected at week 2 p.i. was analyzed. The consensus sequence of nts 298-9375 of the recovered genomes was determined by direct sequencing of PCR products obtained in long RT-PCR followed by nested PCR of 10 overlapping fragments. The identity to clone pCV-H77C sequence was 100%. The consensus sequence of nts 96-291,1328-1848, 3585-4106, 4763-5113 and 9322-9445 was determined from PCR products obtained in different nested RT-PCR assays. The identity of these sequences with pCV-

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H77C was also 100%. These latter regions contained 4 mutations unique to the consensus chimera, including the artificial Afl II cleavage site in the 3' UTR. Therefore, RNA transcripts of this clone of HCV were infectious.

The infectious nature of the consensus chimera indicates that the regions of the 5' and 3' UTRs incorporated into the cassette vector do not destroy viability. This makes it highly advantageous to use the cassette vector to construct infectious cDNA clones of other HCV strains when the consensus sequence for each ORF is inserted.

In addition, two complete full-length clones (dubbed pH21 and pCV-H11) constructed were not infectious, as shown by intrahepatic injection of chimpanzees with the corresponding RNA transcripts. Thus, injection of the transcription mixture into 3 sites of the exposed liver resulted in no observable HCV replication and weekly serum samples were negative for HCV RNA at weeks 1 - 17 p.i. in a highly sensitive nested RT-PCR assay. The cDNA template injected along with the RNA transcripts was also not detected in this assay.

Moreover, the chimpanzee remained negative for antibodies to HCV throughout the follow-up. Subsequent sequence analysis revealed that 7 of 16 additional clones were defective for polyprotein synthesis and all clones had multiple amino acid mutations compared with the consensus sequence of the parent strain. For example, clone pH21, which was not infectious, had 7 amino acid substitutions in the entire predicted polyprotein compared with the consensus sequence of H77 (Fig. 3). The most

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notable mutation was at position 1026, which changed L to Q, altering the cleavage site between NS2 and NS3 (Reed (1995)). Clone pCV-H11, also non-infectious, had 21 amino acid substitutions in the predicted polyprotein compared with the consensus sequence of H77 (Fig. 3). The amino acid mutation at position 564 eliminated a highly conserved C residue in the E2 protein (Okamoto (1992a)).

#### EXAMPLE 4A

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The chimpanzee of Example 4, designated 1530, was monitored out to 32 weeks p.i. for serum enzyme levels (ALT) and the presence of anti-HCV antibodies, HCV RNA, and liver histopathology. The results are shown in Figure 18B.

A second chimp, designated 1494, was also transfected with RNA transcripts of the pCV-H77C clone and monitored out to 17 weeks p.i. for the presence of anti-HCV antibodies, HCV RNA and elevated serum enzyme levels. The results are shown in Figure 18A.

## MATERIALS AND METHODS for Examples 5-10

# 25 Source Of HCV Genotype 1b

An infectious plasma pool (second chimpanzee passage) containing strain HC-J4, genotype 1b, was prepared from acute phase plasma of a chimpanzee experimentally infected with serum containing HC-J4/91 (Okamoto et al., 1992b). The HC-J4/91 sample was obtained from a first chimpanzee passage during the chronic phase of hepatitis C about 8 years after experimental infection.

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The consensus sequence of the entire genome, except for the very 3' end, was determined previously for HC-J4/91 (Okamoto et al., 1992b).

# 5 Preparation Of HCV RNA

Viral RNA was extracted from 100  $\mu$ l aliquots of the HC-J4 plasma pool with the TRIzol system (GIBCO BRL). The RNA pellets were each resuspended in 10  $\mu$ l of 10 mM dithiothreitol (DTT) with 5% (vol/vol) RNasin (20-40 units/ $\mu$ l) (Promega) and stored at -80°C or immediately used for cDNA synthesis.

# Amplification And Cloning Of The 3' UTR

A region spanning from NS5B to the conserved region of the 3' UTR was amplified in nested RT-PCR using the procedure of Yanagi et al., (1997).

In brief, the RNA was denatured at 65°C for 2 minutes, and cDNA was synthesized at 42°C for 1 hour with Superscript II reverse transcriptase (GIBCO BRL) and primer H3'X58R (Table 1) in a 20  $\mu$ l reaction volume. The cDNA mixture was treated with RNase H and RNase T1 (GIBCO BRL) at 37°C for 20 minutes. The first round of PCR was performed on 2  $\mu$ l of the final cDNA mixture in a total volume of 50  $\mu$ l with the Advantage cDNA polymerase mix (Clontech) and external primers H9261F (Table 1) and H3'X58R (Table 1). In the second round of PCR [internal primers H9282F (Table 1) and H3'X45R (Table 1)], 5  $\mu$ l of the first round PCR mixture was added to 45  $\mu$ l of the PCR reaction mixture. Each round of PCR (35 cycles), was

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performed in a DNA thermal cycler 480 (Perkin Elmer) and consisted of denaturation at 94°C for 1 minute (1st cycle: 1 minute 30 sec), annealing at 60°C for 1 minute and elongation at 68°C for 2 minutes. After purification with QIAquick PCR purification kit (QIAGEN), digestion with HindIII and XbaI (Promega), and phenol/chloroform extraction, the amplified products were cloned into pGEM-9zf(-) (Promega) (Yanagi et al., 1997).

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### Amplification And Cloning Of The Entire ORF

A region from within the 5' UTR to the variable region of the 3' UTR of strain HC-J4 was amplified by long RT-PCR (Fig. 1) (Yanagi et al., 1997). The cDNA was synthesized at 42°C for 1 hour in a 20 μl reaction volume with Superscript II reverse transcriptase and primer J4-9405R (5'-GCCTATTGGCCTGGAGTGGTTAGCTC-3'), and treated with RNases as above. The cDNA mixture (2 μl) was amplified by long PCR with the Advantage cDNA polymerase mix and primers A1 (Table 1) (Bukh et al., 1992; Yanagi et al., 1997) and J4-9398R (5'-

25 AGGATGGCCTTAAGGCCTGGAGTGGTTAGCTCCCCGTTCA-3'). Primer J49398R contained extra bases (bold) and an artificial AflII
cleavage site (underlined). A single PCR round was
performed in a Robocycler thermal cycler (Stratagene), and
consisted of denaturation at 99°C for 35 seconds,
annealing at 67°C for 30 seconds and elongation at 68°C
for 10 minutes during the first 5 cycles, 11 minutes
during the next 10 cycles, 12 minutes during the following
10 cycles and 13 minutes during the last 10 cycles.

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After digesting the long PCR products obtained from strain HC-J4 with PinAI (isoschizomer of AgeI) and BfrI (isoschizomer of AflII) (Boehringer Mannheim), attempts were made to clone them directly into a cassette vector (pCV), which contained the 5' and 3' termini of strain H77 (Figure 1) but no full-length clones were obtained. Accordingly, to improve the efficiency of cloning, the PCR product was further digested with BglII (Boehringer Mannheim) and the two resultant genome fragments [L fragment: PinAI/BglII, nts 156 - 8935; S fragment: BglII/BrfI, nts 8936 - 9398] were separately cloned into pCV (Figure 6).

DH5 $\alpha$  competent cells (GIBCO BRL) were transformed and selected on LB agar plates containing 100  $\mu$ g/ml ampicillin (SIGMA) and amplified in LB liquid cultures at 30°C for 18-20 hours.

Sequence analysis of 9 plasmids containing the S fragment (miniprep samples) and 9 plasmids containing the L fragment (maxiprep samples) were performed as described previously (Yanagi et al., 1997). Three L fragments, each encoding a distinct polypeptide, were cloned into pCV-J4S9 (which contained an S fragment encoding the consensus amino acid sequence of HC-J4) to construct three chimeric full-length HCV cDNAs (pCV-J4L2S, pCV-J4L4S and pCV-J4L6S) (Fig. 6). Large scale preparation of each clone was performed as described previously with a QIAGEN plasmid Maxi kit (Yanagi et al., 1997) and the authenticity of each clone was confirmed by sequence analysis.

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### Sequence Analysis

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Both strands of DNA were sequenced with the ABI PRISM Dye Termination Cycle Sequencing Ready Reaction Kit using Taq DNA polymerase (Perkin Elmer) and about 90 specific sense and antisense primers. Analyses of genomic sequences, including multiple sequence alignments and tree analyses, were performed with GeneWorks (Oxford Molecular Group) (Bukh et al., 1995).

The consensus sequence of strain HC-J4 was determined by direct sequencing of PCR products (nts 11 - 9412) and by sequence analysis of multiple cloned L and S fragments (nts 156 -9371). The consensus sequence of the 3' UTR (3' variable region, polypyrimidine tract and the first 16 nucleotides of the conserved region) was determined by analysis of 24 cDNA clones.

# Intrahepatic Transfection Of A Chimpanzee With Transcribed RNA

Two in vitro transcription reactions were performed with each of the three full-length clones. In each reaction 10 μg of plasmid DNA linearized with Xba I (Promega) was transcribed in a 100 μl reaction volume with T7 RNA polymerase (Promega) at 37°C for 2 hours as described previously (Yanagi et al., 1997). Five μl of the final reaction mixture was analyzed by agarose gel electrophoresis and ethidium bromide staining (Fig. 5). Each transcription mixture was diluted with 400 μl of ice-cold phosphate-buffered saline without calcium or magnesium and then the two aliquots from the same cDNA

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clone were combined, immediately frozen on dry ice and stored at -80°C. Within 24 hours after freezing the transcription mixtures were injected into the chimpanzee by percutaneous intrahepatic injection that was guided by ultrasound. Each inoculum was individually injected (5-6 sites) into a separate area of the liver to prevent complementation or recombination. The chimpanzee was maintained under conditions that met all requirements for its use in an approved facility.

Serum samples were collected weekly from the chimpanzee and monitored for liver enzyme levels and anti-HCV antibodies. Weekly samples of 100  $\mu$ l of serum were tested for HCV RNA in a sensitive nested RT-PCR assay (Bukh et al., 1992, Yanagi et al., 1996) with AmpliTaq Gold DNA polymerase. The genome equivalent (GE) titer of HCV was determined by testing 10-fold serial dilutions of the extracted RNA in the RT-PCR assay (Yanagi et al., 1996) with 1 GE defined as the number of HCV genomes present in the highest dilution which was positive in the RT-nested PCR assay.

infectious in vivo, the NS3 region (nts 3659 - 4110) from the chimpanzee serum was amplified in a highly sensitive and specific nested RT-PCR assay with AmpliTaq Gold DNA polymerase and the PCR products were cloned with a TA cloning kit (Invitrogen). In addition, the consensus sequence of the nearly complete genome (nts 11 - 9441) was determined by direct sequencing of overlapping PCR products.

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### EXAMPLE 5

# Sequence Analysis Of Infectious Plasma Pool Of Strain HC-J4 Used As The Cloning Source

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As an infectious cDNA clone of a genotype la strain of HCV had been obtained only after the ORF was engineered to encode the consensus polypeptide (Kolykhalov et al., 1997; Yanagi et al., 1997), a detailed sequence analysis of the cloning source was performed to determine the consensus sequence prior to constructing an infectious cDNA clone of a 1b genotype.

A plasma pool of strain HC-J4 was prepared from acute phase plasmapheresis units collected from a chimpanzee experimentally infected with HC-J4/91 (Okamoto et al., 1992b). This HCV pool had a PCR titer of  $10^4$  -  $10^5$  GE/ml and an infectivity titer of approximately  $10^3$  chimpanzee infectious doses per ml.

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The heterogeneity of the 3' UTR of strain HC-J4 was determined by analyzing 24 clones of nested RT-PCR product. The consensus sequence was identical to that previously published for HC-J4/91 (Okamoto et al., 1992b), except at position 9407 (see below). The variable region consisted of 41 nucleotides (nts. 9372 - 9412), including two in-frame termination codons. Furthermore, its sequence was highly conserved except at positions 9399 (19 A and 5 T clones) and 9407 (17 T and 7 A clones). The poly U-UC region varied slightly in composition and greatly in length (31-162 nucleotides). In the conserved region, the first 16 nucleotides of 22 clones were identical to those previously published for other genotype

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1 strains, whereas two clones each had a single point mutation. These data suggested that the structural organization at the 3' end of HC-J4 was similar to that of the infectious clone of a genotype la strain of Yanagi et al (1997).

Next, the entire ORF of HC-J4 was amplified in a single round of long RT-PCR (Figure 5). The original plan was to clone the resulting PCR products into the *PinAI* and *BrfI* site of a HCV cassette vector (pCV), which had fixed 5' and 3' termini of genotype 1a (Yanagi et al., 1997) but since full-length clones were not obtained, two genome fragments (L and S) derived from the long RT-PCR products (Figure 6) were separately subcloned into pCV.

To determine the consensus sequence of the ORF, the sequence of 9 clones each of the L fragment (pCV-J4L) and of the S fragment (pCV-J4S) was determined and quasispecies were found at 275 nucleotide (3.05 %) and 78 amino acid (2.59 %) positions, scattered throughout the 9030 nts (3010 aa) of the ORF (Figure 7). Of the 161 nucleotide substitutions unique to a single clone, 71% were at the third position of the codon and 72 % were silent.

Each of the nine L clones represented the near complete ORF of an individual genome. The differences among the L clones were 0.30 - 1.53% at the nucleotide and 0.31 - 1.47% at the amino acid level (Figure 8). Two clones, L1 and L7, had a defective ORF due to a single nucleotide deletion and a single nucleotide insertion, respectively. Even though the HC-J4 plasma pool was obtained in the early acute phase, it appeared to contain

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at least three viral species (Figure 9). Species A contained the L1, L2, L6, L8 and L9 clones, species B the L3, L7 and L10 clones and species C the L4 clone. Although each species A clone was unique all A clones differed from all B clones at the same 20 amino acid sites and at these positions, species C had the species A sequence at 14 positions and the species B sequence at 6 positions (Figure 7).

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Okamoto and coworkers (Okamoto et al., 1992b) previously determined the nearly complete genome consensus sequence of strain HC-J4 in acute phase serum of the first chimpanzee passage (HC-J4/83) as well as in chronic phase serum collected 8.2 years later (HC-J4/91). In addition, they determined the sequence of amino acids 379 to 413 (including HVR1) and amino acids 468 to 486 (including HVR2) of multiple individual clones (Okamoto et al., 1992b).

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It was found by the present inventors that the sequences of individual genomes in the plasma pool collected from a chimpanzee inoculated with HC-J4/91 were all more closely related to HC-J4/91 than to HC-J4/83 (Figures 8, 9) and contained HVR amino acid sequences closely related to three of the four viral species previously found in HC-J4/91 (Figure 10).

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Thus, the data presented herein demonstrate the occurrence of the simultaneous transmission of multiple species to a single chimpanzee and clearly illustrates the difficulties in accurately determining the evolution of HCV over time since multiple species with significant changes throughout the HCV genome can be present from the

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onset of the infection. Accordingly, infection of chimpanzees with monoclonal viruses derived from the infectious clones described herein will make it possible to perform more detailed studies of the evolution of HCV in vivo and its importance for viral persistence and pathogenesis.

## EXAMPLE 6

10 Determination Of The Consensus Sequence Of HC-J4 In The Plasma Pool

> The consensus sequence of nucleotides 156-9371 of HC-J4 was determined by two approaches. approach, the consensus sequence was deduced from 9 clones of the long RT-PCR product. In the other approach the long RT-PCR product was reamplified by PCR as overlapping fragments which were sequenced directly. The two "consensus" sequences differed at 31 (0.34%) of 9216 nucleotide positions and at 11 (0.37%) of 3010 deduced amino acid positions (Figure 7). At all of these positions a major quasispecies of strain HC-J4 was found in the plasma pool. At 9 additional amino acid positions the cloned sequences displayed heterogeneity and the direct sequence was ambiguous (Figure 7). Finally, it should be noted that there were multiple amino acid positions at which the consensus sequence obtained by direct sequencing was identical to that obtained by cloning and sequencing even though a major quasispecies was detected (Figure 7).

For positions at which the two "consensus" sequences of HC-J4 differed, both amino acids were

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included in a composite consensus sequence (Figure 7).

However, even with this allowance, none of the 9 L clones analyzed (aa 1 - 2864) had the composite consensus sequence: two clones did not encode the complete polypeptide and the remaining 7 clones differed from the consensus sequence by 3 - 13 amino acids (Figure 7).

## EXAMPLE 7

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Construction Of Chimeric Full-Length cDNA Clones Containing The Entire ORF Of HC-J4

The cassette vector used to clone strain H77 was used to construct an infectious cDNA clone containing the ORF of a second subtype.

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In brief, three full-length cDNA clones were constructed by cloning different L fragments into the PinAI/BglII site of pCV-J4S9, the cassette vector for genotype 1a (Figure 6), which also contained an S fragment encoding the consensus amino acid sequence of HC-J4. Therefore, although the ORF was from strain HC-J4, most of the 5' and 3' terminal sequences originated from strain H77. As a result, the 5' and 3' UTR were chimeras of genotypes 1a and 1b (Figure 11).

The first 155 nucleotides of the 5' UTR were from strain H77 (genotype 1a), and differed from the authentic sequence of HC-J4 (genotype 1b) at nucleotides 11, 12, 13, 34 and 35. In two clones (pCV-J4L2S, pCV-J4L6S) the rest of the 5' UTR had the consensus sequence of HC-J4, whereas the third clone (pCV-J4L4S) had a single nucleotide insertion at position 207. In all 3 clones the first 27 nucleotides of the 3' variable region of the 3'

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UTR were identical with the consensus sequence of HC-J4. The remaining 15 nucleotides of the variable region, the poly U-UC region and the 3' conserved region of the 3' UTR had the same sequence as an infectious clone of strain H77 (Figure 11).

None of the three full-length clones of HC-J4 had the ORF composite consensus sequence (Figures 7, 12). The pCV-J4L6S clone had only three amino acid changes: Q for R at position 231 (E1), V for A at position 937 (NS2) and T for S at position 1215 (NS3). The pCV-J4L4S clone had 7 amino acid changes, including a change at position 450 (E2) that eliminated a highly conserved N-linked glycosylation site (Okamoto et al., 1992a). Finally, the pCV-J4L2S clone had 9 amino acid changes compared with the consensus sequence of HC-J4. A change at position 304 (E1) mutated a highly conserved cysteine residue (Bukh et al., 1993; Okamoto et al., 1992a).

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### EXAMPLE 8

# Transfection Of A Chimpanzee By In Vitro Transcripts Of A Chimeric cDNA

was determined by ultra-sound-guided percutaneous intrahepatic injection into the liver of a chimpanzee of the same amount of cDNA and transcription mixture for each of the clones (Figure 5). This procedure is a less invasive procedure than the laparotomy procedure utilized by Kolykhalov et al. (1997) and Yanagi et al. (1997) and should facilitate in vivo studies of cDNA clones of HCV in

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chimpanzees since percutaneous procedures, unlike laparotomy, can be performed repeatedly.

As shown in Figure 13, the chimpanzee became infected with HCV as measured by increasing titers of  $10^2$  GE/ml at week 1 p.i.,  $10^3$  GE/ml at week 2 p.i. and  $10^4$  -  $10^5$  GE/ml during weeks 3 to 10 p.i.

The viremic pattern found in the early phase of the infection was similar to that observed for the recombinant H77 virus in chimpanzees (Bukh et al., unpublished data; Kolykhalov et al., 1997; Yanagi et al., 1997). The chimpanzee transfected in the present study was chronically infected with hepatitis G virus (HGV/GBV-C) (Bukh et al., 1998) and had a titer of 10<sup>6</sup> GE/ml at the time of HCV transfection. Although HGV/GBV-C was originally believed to be a hepatitis virus, it does not cause hepatitis in chimpanzees (Bukh et al., 1998) and may not replicate in the liver (Laskus et al., 1997). The present study demonstrated that an ongoing infection of HGV/GBV-C did not prevent acute HCV infection in the chimpanzee model.

length HC-J4 clones were infectious, the NS3 region (nts. 3659 - 4110) of HCV genomes amplified by RT-PCR from serum samples taken from the infected chimpanzee during weeks 2 and 4 post-infection (p.i.) were cloned and sequenced. As the PCR primers were a complete match with each of the original three clones, this assay should not have preferentially amplified one virus over another. Sequence analysis of 26 and 24 clones obtained at weeks 2 and 4

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p.i., respectively, demonstrated that all originated from the transcripts of pCV-J4L6S.

Moreover, the consensus sequence of PCR products of the nearly complete genome (nts. 11-9441), amplified from serum obtained during week 2 p.i., was identical to the sequence of pCV-J4L6S and there was no evidence of quasispecies. Thus, RNA transcripts of pCV-J4L6S, but not of pCV-J4L2S or pCV-J4L4S, were infectious in vivo. The data in Figure 13 is therefore the product of the transfection of RNA transcripts of pCV-J4L6S.

In addition, the chimeric sequences of genotypes la and 1b in the UTRs were maintained in the infected chimpanzee. The consensus sequence of nucleotides 11 -341 of the 5' UTR and the variable region of the 3' UTR, amplified from serum obtained during weeks 2 and 4 p.i., had the expected chimeric sequence of genotypes la and lb (Fig. 11). Also three of four clones of the 3' UTR obtained at week 2 p.i. had the chimeric sequence of the variable region, whereas a single substitution was noted in the fourth clone. However, in all four clones the poly U region was longer (2-12 nts) than expected. Also, extra C and G residues were observed in this region. For the most part, the number of C residues in the poly UC region was maintained in all clones although the spacing varied. As shown previously, variations in the number of U residues can reflect artifacts introduced during PCR amplification (Yanagi et al., 1997). The sequence of the first 19 nucleotides of the conserved region was maintained in all four clones. Thus, with the exception of the poly U-UC region, the genomic sequences recovered

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from the infected chimpanzee were exactly those of the chimeric infectious clone pCV-J4BL6S.

The results presented in Figure 13 therefore demonstrate that HCV polypeptide sequences other than the consensus sequence can be infectious and that a chimeric genome containing portions of the H77 termini could produce an infectious virus. In addition, these results showed for the first time that it is possible to make infectious viruses containing 5' and 3' terminal sequences specific for two different subtypes of the same major genotype of HCV.

### EXAMPLE 9

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# Construction Of A Chimeric 1a/1b Infectious Clone

A chimeric 1a/1b infectious clone in which the structural region of the genotype 1b infectious clone is inserted into the 1a clone of Yanagi et al. (1997) is constructed by following the protocol shown in Figure 15. The resultant chimera contains nucleotides 156-2763 of the 1b clone described herein inserted into the 1a clone of Figures 4A-4F. The sequences of the primers shown in Figure 15 which are used in constructing this chimeric clone, designated pH77CV-J4, are presented below.

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- 1. H2751S (Cla I/Nde I)
  CGT CAT CGA TCC TCA GCG GGC ATA TGC ACT GGA CAC GGA
- 2. <u>H2870R</u>
  CAT GCA CCA GCT GAT ATA GCG CTT GTA ATA TG
- 5 3. <u>H7851S</u> TCC GTA GAG GAA GCT TGC AGC CTG ACG CCC
  - 4. <u>H9173 R(P-M)</u>
    GTA CTT GCC ACA TAT AGC AGC CCT GCC TCC TCT G
- 10 5. H9140S (P-M)
  CAG AGG AGG CAG GGC TGC TAT ATG TGG CAA GTA C
  - 6. <u>H9417R</u>
    CGT CTC TAG ACA GGA AAT GGC TTA AGA GGC CGG AGT GTT
    TAC C
- 7.  $\frac{\text{J4-2271S}}{\text{TGC AAT TGG ACT CGA GGA GAG CGC TGT AAC TTG GAG}}$ 
  - 8. <u>J4-2776R (Nde I)</u> CGG TCC AAG GCA TAT GCT CGT GGT AAC GCC AG

20 sequence is shown in Figures 16A-16F) are then produced and transfected into chimpanzees by the methods described in the Materials and Methods section herein and the transfected animals are then be subjected to biochemical

(ALT levels), histopathological and PCR analyses to determine the infectivity of the chimeric clone.

### EXAMPLE 10

# 30 Construction of 3' Deletion Mutants Of The la Infectious Clone pCV-H77C

Seven constructs having various deletions in the 3' untranslated region (UTR) of the 1a infectious clone pCV-H77C were constructed as described in Figures 17A-17B.

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The 3' untranslated sequence remaining in each of the seven constructs following their respective deletions is

shown in Figures 17A-17B.

Construct pCV-H77C(-98X) containing a deletion of the 3'-most 98 nucleotide sequences in the 3'-UTR was transcribed in vitro according to the methods described herein and 1 ml of the diluted transcription mixture was percutaneously transfected into the liver of a chimpanzee with the aid of ultrasound. After three weeks, the transfection was repeated. The chimpanzee was observed to be negative for hepatitis C virus replication as measured by RT-PCR assay for 5 weeks after transfection. results demonstrate that the deleted 98 nucleotide 3'-UTR sequence was critical for production of infectious HCV and appear to contradict the reports of Dash et al. (1996) and Yoo et al. (1995) who reported that RNA transcripts from cDNA clones of HCV-1 and HCV-N lacking the terminal 98 conserved nucleotides at the very 3' end of the 3'-UTR resulted in viral replication after transfection into human hematoma cell lines.

Transcripts of the (-42X) mutant (Figure 17C) were also produced and transfected into a chimpanzee and transcripts of the other five deletion mutants shown in Figures 17D-17G) are to be produced and transfected into chimpanzees by the methods described herein. All transfected animals are to then be assayed for viral replication via RT-PCR.

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### Discussion

In two recent reports on transfection of chimpanzees, only those clones engineered to have the independently determined and slightly different consensus amino acid sequence of the polypeptide of strain H77 were infectious (Kolykhalov et al., 1997; Yanagi et al., 1997). Although the two infectious clones differed at four amino acid positions, these differences were represented in a major component of the quasispecies of the cloning source. In the present study, a single consensus sequence of strain HC-J4 could not be defined because the consensus sequence obtained by two different approaches (direct sequencing and sequencing of cloned products) differed at 20 amino acid positions, even though the same genomic PCR product was analyzed. The infectious clone differed at two positions from the composite amino acid consensus sequence, from the sequence of the 8 additional HC-J4 clones analyzed in this study and from published sequences of earlier passage samples. An additional amino acid differed from the composite consensus sequence but was found in two other HC-J4 clones analyzed in this study. The two non-infectious full-length clones of HC-J4 differed from the composite consensus sequence by only 7 and 9 amino acid differences. However, since these clones had the same termini as the infectious clone (except for a single nucleotide insertion in the 5' UTR of pCV-J4L4S), one or more of these amino acid changes in each clone was apparently deleterious for the virus.

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It was also found in the present study that HC-J4, like other strains of genotype 1b (Kolykhalov et al., 1996; Tanaka et al., 1996; Yamada et al., 1996), had a poly U-UC region followed by a terminal conserved element. The poly U-UC region appears to vary considerably so it was not clear whether changes in this region would have a significant effect on virus replication. On the other hand, the 3'98 nucleotides of the HCV genome were previously shown to be identical among other strains of genotypes 1a and 1b (Kolykhalov et al., 1996; Tanaka et al., 1996). Thus, use of the cassette vector would not alter this region except for addition of 3 nucleotides found in strain H77 between the poly UC region and the 3'98 conserved nucleotides.

In conclusion, an infectious clone representing a genotype 1b strain of HCV has been constructed. Thus, it has been demonstrated that it was possible to obtain an infectious clone of a second strain of HCV. In addition, it has been shown that a consensus amino acid sequence was not absolutely required for infectivity and that chimeras between the UTRs of two different genotypes could be viable.

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WHAT IS CLAIMED IS:

1. A purified and isolated nucleic acid molecule which encodes human hepatitis C virus, said molecule capable of expressing said virus when transfected into cells.

- 2. The nucleic acid molecule of claim 1, wherein said molecule encodes the amino acid sequence shown in Figures 14G-14H.
- 3. The nucleic acid molecule of claim 2, wherein said molecule comprises the nucleic acid sequence shown in Figures 14A-14F.
- 4. The nucleic acid molecule acid molecule of claim 1, wherein said molecule encodes the amino acid sequence shown in Figures 4G-4H.
  - 5. The nucleic acid molecule of claim 4, wherein said molecule comprises the nucleic acid sequence shown in Figures 4A-4F.
  - 6. The nucleic acid molecule of claim 1, wherein a fragment of said molecule which encodes the structural region of hepatitis C virus has been replaced by the structural region from the genome of another hepatitis C virus strain.
  - 7. The nucleic acid molecule of claim 6, wherein said molecule encodes the amino acid sequence shown in Figures 16G-16H.
- 8. The nucleic acid molecule of claim 7, wherein said molecule comprises the nucleic acid sequence shown in Figures 16A-16F.

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- 9. The nucleic acid molecule of claim 1, wherein a fragment of the nucleic acid molecule which encodes at least one HCV protein has been replaced by a fragment of the genome of another hepatitis C virus strain which encodes the corresponding protein.
- 10. The nucleic acid molecule of claim 9, wherein the protein is selected from the group consisting of E1, E2 or NS4 proteins.
- 11. The nucleic acid molecule of claim 1, wherein a fragment of the molecule encoding all or part of an HCV protein has been deleted.
  - 12. The nucleic acid molecule of claim 11, wherein the HCV protein is selected from the group consisting of P7, NS4B or NS5A proteins.
    - 13. A DNA construct comprising a nucleic acid molecule according to claims 1, 3, 5 or 8.
  - 14. An RNA transcript of the DNA construct of claim 13.
    - 15. A cell transfected with the DNA construct of claim 13.
    - 16. A cell transfected with RNA transcript of claim 14.
      - 17. A hepatitis C virus polypeptide produced by the cell of claim 15.
      - 18. A hepatitis C virus polypeptide produced by the cell of claim 16.
- 30 19. A hepatitis C virus produced by the cell of claim 13.
  - 20. A hepatitis C virus produced by the cell of claim 14.

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- 21. A hepatitis C virus whose genome comprises a nucleic acid molecule according to claims 1, 3, 5, 6, 8, or 9.
- 22. A method for producing a hepatitis C virus comprising transfecting a host cell with the RNA transcript of claim 14.
  - 23. A polypeptide encoded by a nucleic acid sequence according to claims 1, 2, 4 or 7 or a fragment thereof.
  - 24. The polypeptide of claim 23, wherein said polypeptide is selected from the group consisting of NS3 protease, El protein, E2 protein or NS4 protein.
  - 25. A method for assaying candidate antiviral agents for activity against HCV, comprising
  - a) exposing a cell containing the hepatitis C virus of claim 21 to the candidate antiviral agent; and
  - b) measuring the presence or absence of hepatitis C virus replication in the cell of step (a).
  - 26. The method of claim 25, wherein said replication in step (b) is measured by at least one of the following: negative strand RT-PCR, quantitative RT-PCR, Western blot, immunofluoresence, or infectivity in a susceptible animal.
  - 27. A method for assaying candidate antiviral agents for activity against HCV, comprising:
    - a) exposing an HCV
       protease encoded by a nucleic acid
       sequence according to claims 1, 2, 4,
       or 7, or a fragment thereof to the

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candidate antiviral agent in the presence of a protease substrate; and b) measuring the protease

activity of said protease.

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The method of claim 27, wherein said HCV 28. protease is selected from the group consisting of an NS3 domain protease, an NS3-NS4A fusion polypeptide, or an NS2-NS3 protease.

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- 29. An antiviral agent identified as having antiviral activity for HCV by the method of claim 25.
- 30. An antiviral agent identified as having antiviral activity for HCV by the method of claim 27.
  - Antibody to the polypeptide of claim 23.

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Antibody to the hepatitis C virus of claim 32.

21.

A method for determining the susceptibility of cells in vitro to support HCV infection, comprising the steps of:

vitro;

growing animal cells in a.

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transfecting into said b. cells the nucleic acid of claim 1; and determining if said

cells show indicia of HCV replication.

The method according to claim 33, wherein said cells are human cells.

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A cassette vector for cloning viral genomes, comprising, inserted therein, the nucleic acid sequence according to claim 2, said vector reading in the

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correct phase for the expression of said inserted sequence and having an active promoter sequence upstream thereof.

- 36. The cassette vector of claim 35, wherein the cassette vector is produced from plasmid pCV.
- 37. The cassette vector of claim 35, wherein the vector also contains one or more expressible marker genes.
- 38. The cassette vector of claim 35, wherein the inserted DNA sequence contains at least one ORF of the HCV genome from any strain.
  - 39. The cassette vector of claim 35, wherein the promoter is a bacterial promoter.
- 40. A composition comprising a polypeptide of claim 23 suspended in a suitable amount of a pharmaceutically acceptable diluent or excipient.
- 41. A method for treating hepatitis C viral infection comprising the administration to a animal in need thereof of a clinically effective amount of the composition of claim 40.
- 42. A composition comprising a nucleic acid molecule of claim 1 suspended in a suitable amount of a pharmaceutically acceptable diluent or excipient.
- 43. A method for treating hepatitis C viral infection comprising the administration to an animal in need thereof of a clinically effective amount of the composition of claim 42.

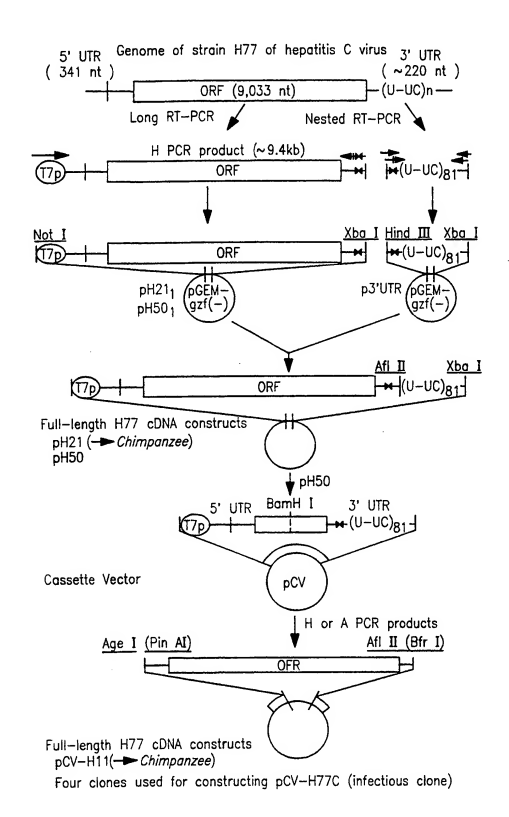
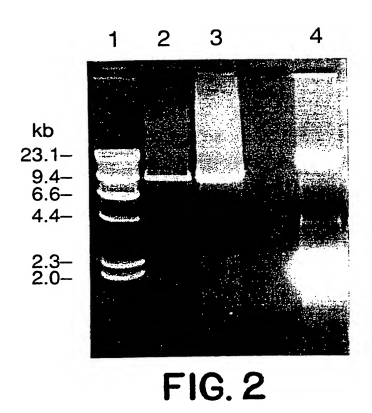
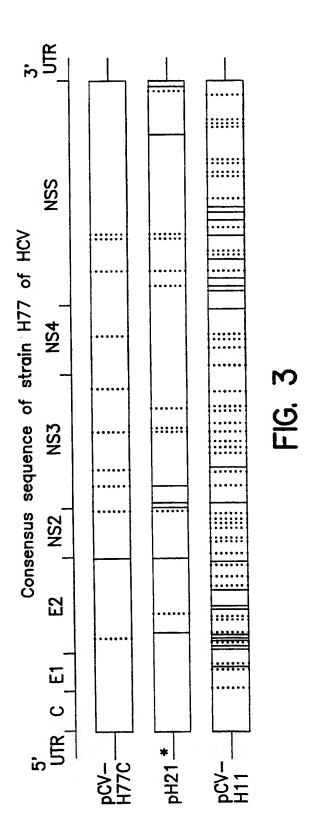


FIG. I





SUBSTITUTE SHEET (RULE 26)

### H77C

	=-	
10 20 30 40	50	
1234567890 1234567890 1234567890 1234567890 1234		
GCCAGCCCCC TGATGGGGGC GACACTCCAC CATGAATCAC TCC		50
GGAACTACTG TCTTCACGCA GAAAGCGTCT AGCCATGGCG TTA		100
TGTCGTGCAG CCTCCAGGAC CCCCCCTCCC GGGAGAGCCA TAG		150
CCCAACCCCT CACTACACCC CAATTCCCAC CACCACCCCC TCC		200
CATAAACCCG CICAAIGCCT GCACATITGG GCGIGCCCCC GCA		250
TAGCCGAGIA GIGIIGGGIC GCGAAAGGCC TIGIGGIACT GCC		300
GICCTICCCA GICCCCCCCC AGGICTCGTA CACCGTCCAC CAT		350
AATCCIAAAC CICAAACAAA AACCAAACGT AACACCAACC GIO		400
GCACGICAAG TICCCGGGIG GCGGICAGAT CGITGGIGGA GIT		450
TGCCGCGCAG GGGCCCTAGA TTGGGTGTGC GCGCGACGAG GAA		500
GAGOGGICGC AACCICGAGG TAGACGICAG CCTATCCCCA AGG		550
GCCCGAGGCC AGGACCIGGG CICAGCCCGG GIACCCIIGG CCC	CICIAIG	600
CCAATGAGGG TIGGGGGIGG GCGGGAIGGC TCCIGICICC CCG	IGGCICT	650
CGCCTACCT GGGGCCCCAC ACACCCCCGG CGTAGGTCGC GCA		700
TAAGGICATO GATACOOTTA OGIGOGGOTT OGCOGACCTO ATG		750
TACCECTOGT CEGCECCCCT CTTGCAGGCG CTGCCAGGGC CCT	GGCGCAT	800
GCCGICCGCG TICIGGAAGA CGCCGIGAAC TAIGCAACAG GGA	ACCITCC	850
TEGITECTET TICTETATET TECTTETESE CETECTETET TEC	CIGACIG	900
TGCCCGCTTC AGCCTACCAA GTGCGCAATT CCTCGGGGCT TTA	CCATGIC	950
ACCAATGATT GCCCIAACIC GAGIATIGIG TACGAGGCGG CCG	ATGCCAT	1000
CCTGCACACT CCGGGGIGIG TCCCTTGCGT TCGCGAGGGT AAC	GCCICGA	1050
CETETTECET CECCETCACC CCCACCETCE CCACCACCECA CCC	CAAACIC	1100
CCCACAACCC AGCITCGACG TCATATCGAT CIGCITGICG GG	£CCCCAC	1150
CCICIGCICG GCCCICIACG TGGGGGACCT GIGGGGGICT GIC	TTTCTTG	1200
TIGGICAACT GITTACCTIC TCTCCCAGGC GCCACTGGAC GAC	CCAAGAC	1250 ~
TGCAATTGIT CTATCTATCC CGCCCATATA ACGCGTCATC GC		1300
CCATATGATG ATGAACTGGT CCCCTACGGC AGCGTTGGTG GT	AGCICAGC	1350
TECTOCECAT COCACAAGOO ATCATEGACA TGATOGOTEG TO	CICACIGG	1400
GCAGICCIGG CGGCATAGC GIATTICICC ATGGIGGGGA AC	IGGGCGAA	1450
GCICCIGGIA GIGCIGCIGC TATTICCOCG CGICCACCCG CA	AACCCACG	1500
TCACCEGEG AAATGCCCEC CECACCACEG CTGGECTTGT TG	GICICCIT	1550
ACACCAGGCG CCAAGCAGAA CATCCAACTG ATCAACACCA AO	GGCAGITG	1600
CCACATCAAT AGCACGGCCT TGAATTGCAA TGAAAGCCTT AA	CACCGGCT	1650
GGITAGCAGG GCTCTTCTAT CAACACAAAT TCAACTCTTC AG	GCIGICCI	1700
CACACCITICG CCACCICCCG ACCCCITIACC CATITICCCC AC	GGCIGGGG	1750
TCCTATCAGT TATGCCAACG GAAGCGCCCT CGACGAACGC CC	CIACIGCI	1800
GCACTACCC TCCAAGACCT TGTGGCATTG TGCCCGCAAA GA	eccicici	1850
GCCCCGGIAT ATTGCTTCAC TCCCAGCCCC GTGGTGGTGG G	ACGACCGA	1900

FIG. 4A

## H77C

10	20	30	40	EA	
1034567890			1234567890	50 1234567890	
			TGCAAATGAT		1950
			GCAATIGGIT		2000
			TGCGGAGCGC		2050
			CIGCCCCACT		2100
GCAAACATCC	GGAAGCCACA	TACICIOGGI	GCCCTCCCC	TCCCTGGATT	2150
ACACCCAGGT	GCATGGTCGA	CIACCOGIAT	AGGCTTTGGC	ACTATOCTEG	2200
TACCATCAAT	TACACCATAT	TCAAAGICAG	GATGIACGIG	GCAGGGGTCG	2250
AGCACAGGCT	GGAAGCGGCC	TGCAACTGGA	CCCCCCCCA	ACCCIGICAT	2300
CIGGAAGACA	GGGACAGGIC	CGAGCTCAGC	CCGITCCICC	TGTCCACCAC	2350
ACAGIGGCAG	GICCITCCGT	GITCTTTCAC	GACCCIGCCA	GCCTTGTCCA	2400
CCCCCCTCAT	CCACCTCCAC	CAGAACATTG	TEGACGIECA	GIACITGIAC	2450
GGGGTAGGGT	CAAGCATCGC	GICCIGGGCC	ATTAAGTGGG	AGTACGTCGT	2500
TCTCCTGTTC	CITCICCITG	CAGAGGGGG	CETCTCCTCC	TGCTTGTGGA	2550
TGATGITACT	CATATCCCAA	GCGGAGGCGG	CTTTGGAGAA	CCICGIAATA	2600
CICAAIGCAG	CATCCCTGGC	CGGGACGCAC	GCTTGTGT	CCTTCCTCGT	2650
GITCITCIGC	TTTGCGTGGT	ATCTGAAGGG	TAGGIGGGIG	CCCCCACCOCC	2700
TCTACGCCCT	CTACGGGATG	TESCETETEC	TCCIGCICCI	GCIGGCGIIG	2750
CCTCAGCGGG	CATACGCACT	GCACACGCAG	GIGGCCCCCI	CCICICCCCC	2800
CGITGITCIT	GICGGGITAA	TGGCGCTGAC	TCIGICGCCA	TATTACAAGC	2850
GCTATATCAG	CIGGIGCATG	TEGIESCITC	AGIATTITCT	GACCAGAGIA	2900
GAAGCGCAAC	TGCACGIGIG	GGLLCCCCCC	CICAACGICC	GGGGGGGG	2950
CGATGCCGTC	ATCTTACTCA	TGIGIGIAGI	ACACCCGACC	CIGGIATTIG	3000
ACATCACCAA	ACTACTCCTG	GCCATCITCG	GACCCCTTTG	GATTCTTCAA	3050
GCCAGTTTGC	TTAAAGICCC	CTACTICGIG	CGCGTTCAAG	GCCLICICCC	3100
			AGGICATTAC		3150
			CCIAIGIGIA		3200
			CTGCGAGATC		
			GACCAAGCTC		
			TCAACGGCIT		
			CCAGCCGACG		
			GGCGIACGCC		
			TGACTGGCCG		3500
			ACTGCTACCC		
			TGICIACCAC		3600
			TCATCCAGAT		3650
			CCICAAGGIT		
			TIACCIGGIC		
CCGATGICAT	TCCCGIGCGC	CGGCGAGGTG	ATAGCAGGG	TAGCCIGCIT	3800

FIG. 4B

# H77C

10	20	30	40	50	
	1234567890	<del>-</del> -	1234567890	1234567890	
	CCATTICCIA				3850
	GGACACGCCG				3900
	TAAAGCGGIG				3950
	CCCCGGIGIT				4000
	CAGGIGGCCC				4050
	CCCGGCTGCG				4100
	CIGITICCICC				4150
	GIIGAICCIA				4200
CIGGCAGCCC	CATCACGIAC	TOCACCTACG	CCAAGITCCT	TGCCGACGGC	4250
GGGIGCTCAG	GAGGIGCITA	TGACATAATA	ATTTGTGACG	AGIGCCACIC	4300
CACGGATGCC	ACATCCATCT	TEGECATOGG	CACIGICCIT	GACCAAGCAG	4350
AGACTGCGGG	GCCCAGACTG	GIIGIGCICG	CCACIGCIAC	CCCICCGGGC	4400
TCCGTCACTG	TGTCCCATCC	TAACATCGAG	CACCITICCIC	TGTCCACCAC	4450
CGGAGAGATC	CCCTTTIACG	GCAAGGCTAT	CCCCCICCAG	GIGATCAAGG	4500
GGGGAAGACA	TCTCATCTTC	TGCCACTCAA	AGAAGAAGIG	CGACGAGCTC	4550
GCCGCGAAGC	TGGTCGCATT	GOGCATCAAT	CCCLCCCL	ACTACCGCGG	4600
TCTTGACGIC	TCIGICATCO	CGACCAGCGC	CGATGITGIC	GICGIGICGA	4650
CCGATGCTCT	CATGACTOGO	TTTACCGGCG	ACTICGACIO	TGTGATAGAC	4700
TGCAACACGI	GIGICACICA	A GACAGICGAI	TICAGCCITC	ACCCIACCIT	4750
TACCATTGAC	ACAACCACG	TCCCCCAGG	r recrerence	AGGACTCAAC	4800
GCCGGGGCA	GACTOGCAG	GGGAAGCCAC	GCATCIATAC	ATTIGIGGCA	4850
CCGGGGGAG	C GCCCCICCCC	G CATGITICGAL	TOGRAGIA	TCTGTGAGTG	4900
CTATGACGC	G GGCTGTGCT	r ggraigagc	r caccoccc	CAGACTACAG	4950
TTAGGCTAC	G AGCGIACAT	G AACACCCCC	GCTTCCCG	GIGCCAGGAC	5000
CATCTIGAA	TTTGGGAGG	G CGICTTIAC	G GCCTCACTO	ATATACATCC	5050
CCACTTTTT	A TOCCAGACA	a accagagic	G GGAGAACIT	r certaceteg	5100
TAGOGTACO	A AGCCACOGI	G TOCGCTAGG	G CTCAAGOOO	TCCCCCATCG	5150
TGGGACCAG	a TGIGGAAGI	G TITGATCOS	C CITAAACCC	A CCCTCCATGG	5200
GCCAACACC	C CIGCIATAC	A GACTGGGGG	C TGITCAGAA	r GAAGICACCC	5250 5300
TGACGCACC	C AATCACCAA	A TACATCAIG	A CATGCATGI	C GGCCGACCIG	5350 5350
GAGGICGIC	'A CGAGCACCI	G GGIGCICGI	T GGCGGGIU	C TGGCTGCTCT	5400
GCCCCCIV	T TECCIGIC	A CAGGCIGOS	T GGICALAGI	G GGCAGGATCG	5450
TCTTGTCCC	GAAGCCGGC	A ATTATACCI	G ACAGGIAGG	T TCTCTACCAG	5500
GAGTICGAT	IG AGATGGAAC	A GIGCICICA	G CACTTACCO	T ACATCGAGCA	5550 5550
AGGGATGA	IG CICCCICAC	C AGITCAAGC	A GAAGGCCI	C GCCICCIGC	5600
AGACCGCG.	IC CCGCCAIG	A GAGGITATO	A CCCIGITO	T CCAGACCAAC	5650
TGGCAGAA	AC TOGAGGIC	TT TIGGGCGAV	AC CALAIGIG	ATTICATCAG	5700
TGGGATAC	AA TACTTGGC	3G GCCIGICA		ET AACCCCGCCA	3,00

FIG. 4C

WO 99/04008

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## H77C

PCT/US98/14688

10	20	30	40	50	
1234567890		1234567890			-
TIGCITCATT		ACAGCTGCCG			5750
GGCCAAACCC	TCCTCTTCAA	CATATIGGG	GGGIGGGIGG	CIGCCCAGCT	5800
accecacac	GGIGCCGCIA	CIGCCTTIGT	GGGIGCIGGC	CTACCTCCCC	5850
CCGCCATCGG	CAGCGITGGA	CIGGGGAAGG	TOCTOGTOGA	CATTCTTGCA	5900
GGGIATGGCG	COCCUTAGE	GGGAGCICIT	GIAGCATICA	AGATCATGAG	5950
CEGICAGGIC	CCCTCCACCG	AGGACCIGGI	CAATCIGCIG	CCCCATCC	6000
TCTCGCCTGG	AGCCCTTGIA	GIOGGIGICG	TCTCCCCAGC	AATACTGCGC	6050
CGGCACGITG	GCCCGGGGGA	GGGGCAGIG	CAATGGATGA	ACCGCTAAT	6100
AGCCTTCGCC	TCCCGGGGGA	ACCATGITTC	CCCCACGCAC	TACGIGOOGG	6150
AGAGCGATGC	AGCCGCCCCC	GICACIGOCA	TACTCAGCAG	CCTCACTGIA	6200
ACCCAGCTCC	TGAGGCGACT	GCATCAGTGG	ATAAGCTCGG	AGIGIACCAC	6250
TOCATGCTCC	GGITCCIGGC	TAAGGGACAT	CIGGGACIGG	ATATGCGAGG	6300
TGCTGAGCGA	CITTAAGACC	TGGCTGAAAG	CCAAGCTCAT	GCCACAACTG	6350
CCTGGGATTC	CCTTTGTGTC	CTGCCAGCGC	GGGTATAGGG	GGGICIGGGG	6400
AGGAGACGGC	ATTATGCACA	CTCGCTGCCA	CIGIGGAGCT	GAGATCACTG	6900
GACATGICAA	AAACGGGACG	ATGAGGATCG	TOGGTOCTAG	CACCTCCACG	6950
AACATGTGGA	GIGGGACGIT	CCCCATTAAC	GCCTACACCA	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	6550
TACICCCCIT	CCIGCGCCGA	ACTATAAGIT	CCCCCIGICG	AGGGIGICIG	6600
CAGAGGAATA	CGIGGAGATA	AGGCGGGTGG	GGGACTICCA	CIACGIAICG	6650
GGIATGACIA	CIGACAAICI	TAAATGCCCG	TECCAGATCC	CATCGCCCGA	6700
ATTITICACA	GAATTGGACG	GGGIGGGCT	ACACAGGITT	GCGCCCCTT	6750
GCAAGCCCTT	GCIGCGGGAG	GAGGIAICAT	TCAGAGTAGG	ACTCCACGAG	6800
TACCCGGIGG	GGICGCAATT	ACCTTGCGAG	CCCGAACCGG	ACGIAGCOGI	6850
GIIGACGICC	ATGCTCACTG	ATCCCTCCCA	TATAACAGCA	CACCCCCCC	6900
GGAGAAGGIT	GGCGAGAGGG	TCACCCCCTT	CTATGGCCAG	CICCICGGCT	6950
AGCCAGCIGI	CCGCTCCATC	TCTCAAGGCA	ACTIGCACCG	CCAACCATGA	7000
	GCCGAGCTCA				7050
	CATCACCAGG				7100
	ATCCGCTIGT				7150
				GCCIGCCC	
	GCCGGACTAC				7250
	AACCACCIGI				7300
	, GIGCCICCGC				7350
	ATCIACIGCC				7400
	CITCOGGCAT				7450
	TCIGGCIGCC				7500
	CCTGGAGGGG			•	7550 7600
TCATGGTCGA	CGGICAGIAG	TGGGGCCGAC	ACGGAAGAIG	TCGIGIGCIG	7600

FIG. 4D

## H77C

	20 30		50	
1234567890 12345678	90 1234567890	1234567890	1234567890	
CICAATGICT TATTCCIG	GA CAGGCGCAC	r cgrcacccg	TECECTECEG	7650
AAGAACAAAA ACTGCCCA				7700
CACAATCIGG TGIATTOO	AC CACTICACO	C AGIGCITGCC	AAAGGCAGAA	7750
CAAAGICACA TITGACAG				7800
TOCTCAAGGA GOTCAAAG				7850
TOOGIAGAGG AAGCITGO	AG CCTGACGCO	CCACATICAG	CCAAATCCAA	7900
GITTGGCTAT GGGGCAAA	AG ACGICCGITI	G CCATCCCAGA	AAGGCCGTAG	7950
CCCACATCAA CTCCGIGI	GG AAAGACCTT	C TGGAAGACAG	TGTAACACCA	8000
ATAGACACTA CCATCATO	CAAGAACGA	G GITTICIGOG	TICAGCCIGA	8050
GAAGGGGGT CGTAAGCC				8100
TECECGIGIG CGAGAAG	ATG GCCCTGTAC	G ACGIGGITAG	CAAGCTCCCC	8150
CTGCCCGTGA TGGGAAG	TIC CTACGGATT	C CAATACTCAC	CAGGACAGCG	8200
GTTGAATTC CTCGTGC	AAG CGIGGAAGI	C CAAGAAGACC	CCGATGGGGT	8250
TCTCGIATGA TACCCGC				8300
CGTACGGAGG AGGCAAT	TTA CCAATGITG	T GACCTGGACC	CCCAAGCCCG	8350
CGTGGCCATC AAGTCCC	TCA CTGAGAGGC	T TIAIGITGG	GCCCICTIA	8400
CCAATTCAAG GGGGGAA	AAC TGCGGCTAC	C GCAGGIGCCC	CCCCACCCCC	8450
GIACIGACAA CIACCIG	TGG TAACACCC	C ACTIGCIACA	A TCAAGGCCCG	8500
GCAGCCTGT CGAGCCG	CAG GGCTCCAG	A CIGCACCAIC	e creerere	8550
GCCACCACIT AGICGIT				8600
COCACCTCA CACCCTT	TAC GGAGGCIA	IG ACCAGGIAC	r accecece	8650
GGGGACCCC CCACAAC	CAG AATACGAC	TT GGAGCITATA	A ACATCATECT	8700
CCTCCAACGT GTCAGTC	CACGACGG	CTCGAAAGA	G GGTCTACTAC	8750
CTTACCCGTG ACCCTAC	AAC CCCCTCG	OG AGAGCCGCG	r gggagacagc	8800
	ATT CCTGGCTA			8850
CCACACTGTG GGCGAG				8900
ATAGCCAGG ATCAGC	TIGA ACAGGCIC	TT AACTGTGAG	A TCTACGGAGC	8950
CIGCIACICC ATAGAAC	CAC TGGATCIA	CC TCCAATCAT	T CAAAGACTCC	9000
ATGGCCTCAG CGCATT	MICA CICCACAC	TT ACTOTOCAG	G TGAAAICAAT	9050
AGGGTGGCCG CATGCC	ICAG AAAACTIC	EG GICCCECCC	T TECCACCTIC	9100
GAGACACCGG GCCCGG	AGCG TCCGCGCT	AG GCTTCTGTC	C AGAGGAGGCA	9150
GGGCTGCCAT ATGTGG	CAAG TACCICIT	CA ACTGGGCAG	T AAGAACAAAG	9200
CICAAACICA CICCAA	TAGC GGCCGCTC	CCCCCCCCAC	T TGTCCGGTTG	9250
GITCACGGCT GGCTAC	AGCG GGGGAGA	AT TTATCACAC	C GIGICICAIG	9300
cccecccc creer	CIGG TITIGCC	PAC TOCTOCTO	C TGCAGGGGIA	9350
GCATCTACC TOCTOC	CCAA CCGATGA	AGG TIGGGGTA	A CACTOOGGCC	9400
TCTPAAGCCA TITCCT	GITT TITTIT	er tereter	TTTTTCTTT	9450
TTTTTTCTT TCCTT	CCTT CTTTTT	ICC TITCITIT	C CCTTCTTTAA	9500
	- <del>-</del> -			

FIG. 4E

# H77C

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
TEGTESCICC	ATCTTAGCCC	TAGTCACGGC	TAGCIGIGAA	AGGICCGIGA	9550
GCCGCATGAC	TGCAGAGAGT	GCTGATACTG	GCCICICIGC	AGATCATGT	9599

FIG. 4F

# H77C

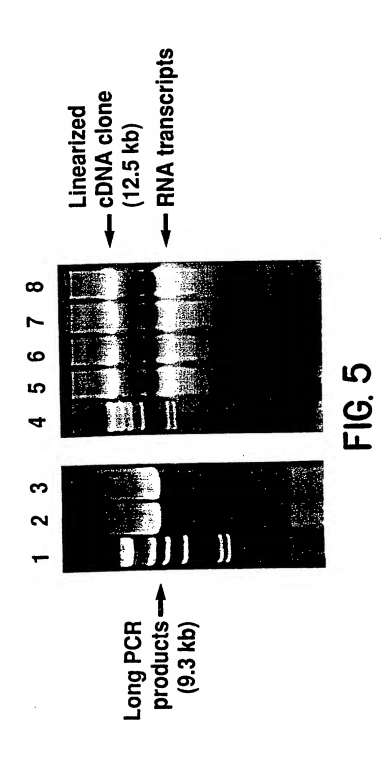
20 40 50	
10 20 30 40 50	
1234567890 1234567890 1234567890 1234567890 1234567890	50
MSINPKPORK IKRNINRRPO DVKFPGGGQI VGGVYLLPRR GPRLGVRAIR	100
KISERSOPRG RROPIPKARR PEGRIWAQPG YPWPLYGNEG CGWAGWLLSP	
RGSRPSWGPT DPRRRSRNLG KVIDTLITCGF ADLMGYIPLV GAPLGGAARA	<b>15</b> 0
LAHGVRVLED GVNYATGNLP GCSFSIFILA LLSCLTVPAS AYQVRNSSGL	200
YHVINDCENS SIVYEAADAI LHIPGCVPCV REGNASRCW AVIPIVATRD	250
CKLPTTOLRR HIDLINGSAT LCSALYVEDL CGSVFLVGQL FTFSPRRHWT	300
TODONOSIYP CHITCHRMAW DMMNWSPIA ALVVAQLLRI PQAIMIMIAG	350
AHWGVIAGIA YFSMVGWAK VLVVILLFAG VDAEIHVIGG NAGRTIAGIV	400
GLLTPGAKON IOLININGSW HINSTALNON ESLNIGWLAG LFYOHKFNSS	450
COPERIASOR RUIDEACGWG PISYANGSGL DERPYCWHYP PRPCGIVPAK	500
SVCGPVYCFT PSPVVVGTID RSCAPIYSWG ANDIDVFVLN NIRPPLGWF	550
COTWINISTIGE TKVCGAPPCV IGGVGNNTLL CPIDCFRKHP EATYSRCGSG	600
PWITTPROMID YPYRLWHYPC TINYTIFKVR MYVGGVEHRL EAAOWIRGE	650
RCDLEDRORS ELSPLILSTY OWOVLPCSFT TLPALSTGLI HLHQNIVDVQ	700
VIVENESSIA SWAIKWEYVV LLFLLLADAR VCSCLWMMLL ISQAFAALEN	750
INTINAASTA GIHGLVSFLV FFCFAWYLKG RWPGAVYAL YGMMPLLLLL	800
LALPORAYAL DIEVAASCGG WINGIMALT LSPYYKRYIS WOMWILQYFL	850
TRY FAOT HIM VPPI NVROGR DAVILLMOVV HPTLVFDITK LLLATFGPLW	900
IIQASILKVP YFVRVQGILR ICALARKIAG GHYVQMAIIK LGALIGIYVY	950
NHLTPLRDWA HNGLRDLAVA VEPVVFSRME TKLTIWGADI AACGDIINGL	1000
EVENERGOFT LIGPADOMVS KOWRLLAPIT AYAQQIRGLL GCIITSLIGR	1050
DENOTIFICATION TO TATATOTET, ATCINGUCWI VYHGAGIRTI ASPKGPVIQM	1100
YINVDODLVG WPAPOGSRSL TPCICGSSDL YLVIRHADVI PVRRGDSRG	1150
SLISPRPISY LKGSSGGPLL CPACHAVGLF RAAVCIRGVA KAVDFIPVEN	1200
LGTIMRSPVF TENSSPPAVP QSFQVAHLHA PIGSGKSTKV PAAYAAQGYK	1250
VLVLNPSVAA TLGFGAYMSK AHGVDPNIRT GVRITTIGSP ITYSTYGKFL	1300
ADGGCSGGAY DILICDECHS TDATSILGIG TVLDQAETAG ARLWLATAT	1350
PPGSVIVSHP NIEEVALSTT GEIPFYGKAI PLEVIKOGRH LIFCHSKKKC	1400
DELAAKLVAL GINAVAYYRG LDVSVIPISG DVVVVSIDAL MIGFIGDFDS	1450
VIDONICVIQ TVDFSLDPIF TIETTILPQD AVSRIQRRGR TGRGKPGIYR	1500
FVAPGERPSG MFDSSVLCEC YDAGCAWYEL TPAETIVRLR AYMNTPGLPV	1550
CODHLEFWEG VFIGLIHIDA HFLSQIKQSG ENFPYLVAYQ AIVCARAQAP	1600
PPSWDOMKC LIRLKPILIG PIPLLYRLGA VONEVILIHP ITKYIMIOMS	1650
ADLEWISIW VLVGGVLAAL AAYCLSIGCV VIVCRIVLSG KPAIIPDREV	1700
LYQEFDEMEE CSQHLPYIEQ GMMLAEQFKQ KALGLLQTAS RHAEVITPAV	1750
QINWOKLEVF WAKHMWAFIS GIQYLAGIST LPGNPAIASL MAFTAAVISP	1800
UTTEQTILEN ILOGWAAQL AAPGAATAFV GAGLAGAAIG SVGLGKVLVD	1850
ILAGYGAGVA GALVAFKIMS GEVPSTEDLV NLLPAILSPG ALVVGVVCAA	1900
TIAGYCAGVA GALVARATIO GENESTEDIN TATTETO INTO	

FIG. 4G

# H77C

10	20	30	40	50	
1234567890	1234567890	123456/890	1234567890	1234307630	1950
II_RRHVGPGE	GAVQWMNRLL	AFASRENHVS	PIHYVPESDA	AARVIALLSS	
LIVIQLLRRL	HOWISSECIT	PCSGSWLRDI	WDWICEVLSD	FKIWLKAKIM	2000
POLPGIPFVS	CORGYRGWR	GDGIMHIRCH	CGAELIGHVK	NGIMRIVGPR	2050
			ALWRVSAEEY		2100
YVSGMITINL	KCPCQIPSPE	FFTELDGVRL	HRFAPPCKPL	LREEVSFRVG	2150
LHEYPVGSQL	PCEPEPDVAV	LISMLIDPSH	ITAFAAGRRL	ARGSPPSMAS	2200
SSASQLSAPS	LKATCIANHD	SPDAELIEAN	LLWRQEMGGN	ITRVESENKV	2250
VILDSFDPLV	<b>AEEDEREVSV</b>	PAEII RKSRR	FARALPWAR	PDYNPPLVET	2300
<b>WKKPDYEPPV</b>	VHGCPLPPPR	SPPVPPPRKK	RIVVLTESIL	STALAELATK	2350
SFGSSSTSGI	TGDVITISSE	PAPSGCPPDS	DVESYSSMPP	LEGEPGDPDL	2400
SDGSWSTVSS	<b>GADIEDWCC</b>	SMSYSWIGAL	VIPCAAEEQK	LPINALSNSL	2450
LRHHNLVYST	TSRSACQRQK	KVIFDRLQVL	DSHYQDVLKE	VKAAASKVKA	2500
NLLSVEEACS	LITPPHSAKSK	FGYGAKDVRC	HARKAVAHIN	SWKDLLEDS	2550
VIPIDITIMA	KNEVFCVOPE	KGGRKPARLI	VFPDLGVRVC	EKMALYDVVS	2600
KT PLAVMGSS	YGFOYSPGOR	VEFLVQAWKS	KKIPMGFSYD	TRCFDSIVIE	2650
SDIRTEFATY	OCCDLDPOAR	VAIKSLIERL	YVGGPLINSR	GENCGYRRCR	2700
ASCAL TABLE	NTLICYIKAR	AACRAAGLOD	CIMLVCGDDL	VVICESAGVQ	2750
			ELITSCSSNV		2800
			NIIMFAPILW		2850
STARME	ONINCETYGA	CYSIEPLDLP	PIIQRLHGLS	AFSLHSYSPG	2900
ETVEX VA VCI D	KT CAPPI RAW	RHRARSVRAR	LLSRGGRAAI	CCKYLFNWAV	2950
עדמיי זען דעיייטרא	AACRI DI GGA	FTAGYSGEDI	YHSVSHARPR	WFWFCILLLA	3000
AGVGTYLLEN					3011
	K		ALL		

FIG. 4H



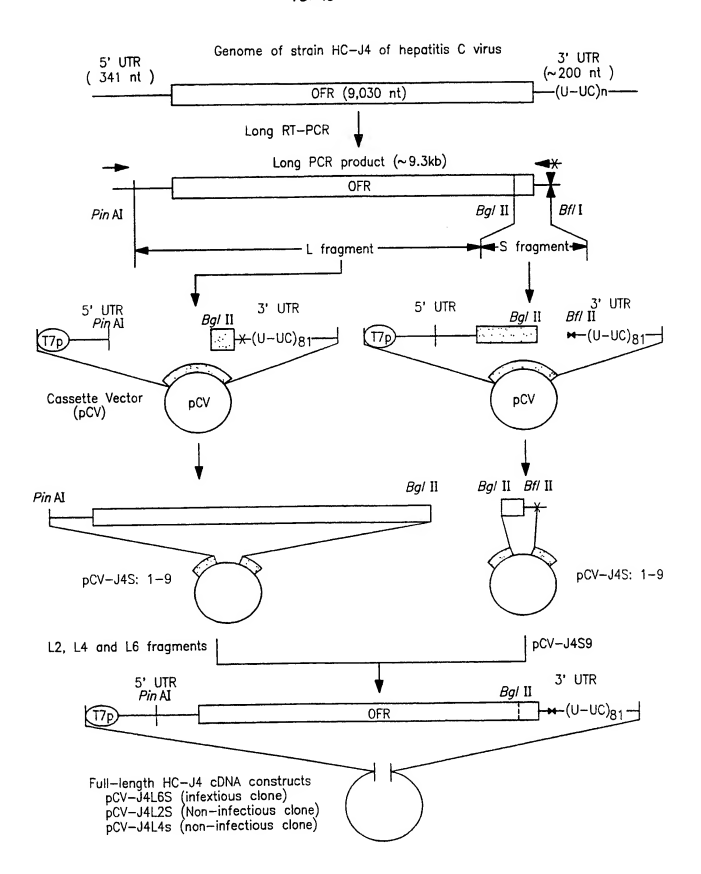


FIG. 6
SUBSTITUTE SHEET (RULE 26)

	_	_	Ť		T	Т	_		<b>T</b>	_	Г	7		1	7	_	T	_	_	٦
Cons-F	2			<b>∠</b> ∀		אים	∢	٥	7	∝	c	و	2		2	L	,	ر	٧	
Cons-D	۰		٠	-		K,U	•			•		۰	•		•	•		۰	۰	
L4(C)	٠	ì	۰		-	•	•	ľ	,	۰		•	,	,	n	٠		•	٠	
L10(B)	,	,		-	-	3	•	-	r	۰		∢	ء		۰	l.		۰	-	
(17*(B)		°	٠	٢	-	0			•	Į,	·	∢	ے	2	۰	Š		٠		
L3(B)		۰	٠	-	-	o	•		Ι		,	⋖	٥	a	۰		,	٥	•	_
(A)61		٥	0		٥	•	,	Ì	٠		٥	۰		۰	•		A	۰		
(A)8(A)			۵		۰	6		°	•	ļ	צכ			٠	۰		•			
16(A)	2	•			•	•		۰	۰	ļ	<b>-</b>	۰		٥	۰		٠	•		•
12(A)	1	•	,	,	•	۰		۰	۰		<b>3</b>	٠		•	۰		٠	>	-	٠
(A)	22	v	À	•	۰	۰		٥	۰		•	,	,	۰	,	ļ	<		°	•
00-000	cd silos	2	-	_	<	۵	2	×	٥	2	~	:   c	و	z	1	2	4	راد	اد	A
framount	ר ונמלווופוור	46	01	36	52	5 5	2/	189	405	65	231	220	722	234	010	nc7	200	202	204	379
			Core			-														

FIG. 7A

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Cons-F	E,T	H,Y	L'S	R,G	>	A,V	Ŧ	S	O,H	7,	A,T	S	S	A,V	>-	K,E	V,I	>	>	_	7	O	A
Cons-D	E,T	H,Y	1,5	G	•	۸	•	•	Ŧ		_	•	•	A,V	•	Ш	I	•	•	•	•	•	•
L4(C)	•	•	•	٠	•	^	œ	•			1	Р	•		•	ш		•	•	•	•	•	•
L10(B)	-	>	S	ပ	A	•	~	٠	Ξ		_	•	•	۸	•	w		٠	•	•	•	•	•
(8) (7)	T	>	S	၁		•	æ	ď	H	1	_	•	٠	۸	•	ш	-	A		>	•	•	>
(a)£1	-	>	S	ပ	•	•	R	•	Τ	Ţ	L	•	٠	۸	Н	E		•	٠	•	•	٠	•
(A)61			•	•		>	٠	•	•	•	•	•	•	•	•	•	•	A	•	٠	۸	•	•
L8(A)		•			•	>	•	٠		•	•	•	z		•	•	•	•	•	•	•	Ь	•
L6(A)		•						•		•	•	•	•		•	•	•	•	•	•	•	•	•
L2(A)						•						•			•		•	•	M	٠	•	•	•
(A) *L1						>	•	•					•		•		•	•	•	•	٨	•	•
Cons-p9		, =	-	. 2	· />	A	H	S	0	L	A	S	S	A	>	×	>	>	>	-		o	A
L fragment	384	386	388	390	391	392	394	405	434	438	444	450	458	466	474	476	496	524	536	580	622	673	783
	5	בל בל	1	1	•	<u>.</u>						•			···								/d

FIG. 7B

16/49

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Cons-F	9	>	7	4>	> <	τ .	A,D	۵	V		1,7	Ľ,'D	-	ى		0	4	A.	4	2	د	-	<b>&gt;</b>	-	-	_
Cons-D	•	•			۰	٥	۵	۰	•	, -	_	H,	0	•		0	•	0			0	۰	•	·		•
L4(C)	·	,		•		٠	۰	1		•	•	•	۰	٠	,	•	•			•	۰	٠	Н			•
L10(B)				•	•		Q		,  -	- -	-	Ŧ	۰		•	٠	•	>	>	•	٠	S	۰		•	•
(B) <sub>*</sub> (7)	U	7		×	۰	•	6			•	-	Ŧ	۰		•		٠				٠	٠				
L3(B)	,			٠	•	۰	ے	,	•	•		エ	•			٠	۰			•	•	S				
(A)61			•	٥		۰		·		٠	۰		۰			۰	۰		•	•	۰	۰				
L8(A)		,		•	I				^	۰	•	٠	۰		•	٠					٠			.	•	
L6(A)			0	•		>				۰	<b></b>	٠	•			<b> </b>	ľ	•		•	•		,	•	٠	٠
L2(A)	- 1	•		۰				·	٠	•	۰	۰			٠	۰	ر				٠			۰	٠	а.
11* (A)			•	۰			,	٠	٠	۰	۰	•	>	<	~	۰		•	•	^	Z			٠	A	۰
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framont	ר ווסליווכיוו	820	857	200	934	- 250	93/	978	1028	1031	1043	1067	200	1097	1188	1915	217	1223	1226	1339	1300	1503	COCI	1528	1535	1662
		NC2	70.	<u>L</u>			1		NC 3	2	<u>.</u>	_1	<del></del>			-				•	•					NS4A

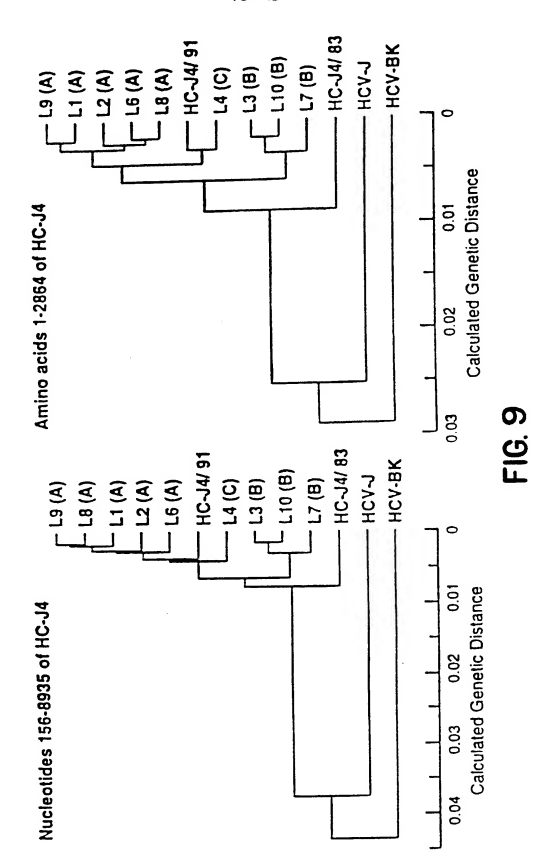
FIG. 70

0	Cons-p9	L1* (A)	L2(A)	L6(A)	L8(A)	(A)G.	(3(8)	(8),	L10(B)	L4(C)	Cons-D	Cons-F K
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		•	•	•	•	•	LL	•	•	•	•	
w		•	•	•	•	•	۵	۵	۵	•	E,D	E,U
>		•	•	•	•	•	П	•	•	•	•	> .
		•	•	•	٠	•	0	0	0		L,0	1,0
>-		•	•	•	•	•	•		•	I	•	<b>-</b>  :
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A		•	•	•	•	•	•	$\cdot  $	•	•	•	W (
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A		•	•	•	٠		>	•	•	•	•	¥
$\bigg/ \bigg/$	1	SS	83	ZS	53	S7	88	S10	\$4	9S		
ပ	1			•	٠	•	S	S	٠	•	•	၁
S		•		•	•	•	9	S	9	ပ	•	S
Ω		•	•	·	•	•		$\cdot$	•	ပ	•	a
S		٠	L.	<u></u>	L.			•		•	•	S

FIG. 70

1 (A)	12 (A)	L6 (A)	L8 (A)	(A) 6J	(8)	(8)	L10 (B)	L4 (C)	HC-J4/91HC-J4/83	HC-34/83
	0.56	0 60	0.36	0.33	1.50	1.53	1.46	0.95	0.83	1.79
$\overline{}$		0.55	0.35	0.50	1.49	1.51	1.45	0.98	0.82	1.77
	040		0.31	0.55	1.33	1.38	1.29	0.80	0.68	1.58
	0.38	0.31		0.31	1.32	1.34	1.28	0.79	0.65	1.62
	0.52	0.45	0.35		1.42	1.42	1.38	0.91	0.75	1.66
	143	1.15	1.33	1.36		0.61	0.30	1.43	0.90	1.51
	1.33	1.05	1.22	1.22	99.0		0.57	1.47	0.95	1.54
1	1.33	0.59	1.22	1.26	0.31	0.56		1.37	0.85	1.42
)	0.80	0.59	0.63	1.26	1.12	1.08	1.01		0.76	1.73
- 1	0.91	0.63	0.80	0.87	0.77	0.73	99.0	0.52		1.22
- 1	1.89	1.68	1.85	1.82	1.75	1.61	1.61	1.71	1.40	
- 1										

F1G, 8



SUBSTITUTE SHEET (RULE 26)

	20/49		
486 YTKPNSS DQRPYC E	ы ы ы 	H. E	ESG.R E.D.P
468 GWGPIT	<b>4</b>		• • •
413 HC-J4L6 (A): AGVDG ETHTTGRVAGHTTSGFTSLFSSGAS QKIQL HC-J4L2 (A): HC-J4/91-20: HC-J4L1 (A): HC-J4L8 (A): HC-J4L9 (A): HC-J4L9 (A):	HC-J4L4 (C):	HC-J4L7 (B) : T.Y.S.G.R. P. HC-J4L10 (B) : T. T.Y.S.GA.R. HC-J4L3 (B) : T. T.Y.S.G.R. HC-J4/91-26 : T. T.Y.S.G.R. HC-J4/91-25 : T.Y.S.G.R. HC-J4/91-25 : A.Y.S.G.R. HC-J4/91-24 : A.Y.S.G.R. HC-J4/91-24 : A.Y.S.G.R. HC-J4/91-24 : A.Y.S.G.R.	HC-J4/91-27: K.Y.S.GA.SRPR HC-J4/83:X.S.GA.STLAPR HVR1 FIG. 10

90 CCC GATIGGGGC GACACICCAC CAIAGAICAC ICCCCIGIGA GGAACIACIG ICTICACGCA GAAAGCGICI AGCCAIGGCG IGA	TGTCGTGCAG CCTCCAGGAC CCCCCTCCC GGGAGAGCCA TAGTGGTCTG CGGA <u>ACCGGT</u> GAGTACACCG GAATTGC	TCCTITCTIG GAICAACCCG CTCAATGCCT GGAGAITIGG GCGTGCCCCC GCGAGACTGC TAGCCGAGTA GTGTTGG	341 271 GCGAAAGGCC TIGIGGIACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGIGGTACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGIGGTACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCCTTGCGA GTGCCCCGGG AGGTCTCGTA GACCGTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCCTTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCCTTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCCTGATAGG GTGCTTGCAC C 6. CCGAAAGGCC TIGITGGTACT GCCTGATAGG GTGCCTGATAGG GTGCTGATAGG GTGCTGATAG			9372 :TGAACGGGGA GCTAACCACT CCAGGCCTTC CTG poly (U-UC) <sub>n</sub> GGTGGCT CCATCTTAG :TGAACGGGGA GCTAACCACT CCAGCCTTC CTG poly (U-UC) <sub>81</sub> AAT :G.TT.G.G.ABfr 1	3' conserved region (Cont.)	GGCTAGCTGT	
; Untranslated Region  1 1c-J4 :GCCAGCCCCC GATTGGGGGC GACV-J4L6S:TGA	91 iC-J4 :TTAGTATGAG TGTCGTGCAG CC oCV-J4L6S:	#C-J4 :GACGACCGGG TCCTTTCTTG G #PCV-J4L6S:	HC-J4 pCV-J4L6S pCV-H77C	3' Untranslated Region	•	HC-J4 pCV-J4L6S pCV-H77C		9514 :CCCTAGTCAC GGCTAGCTGT	

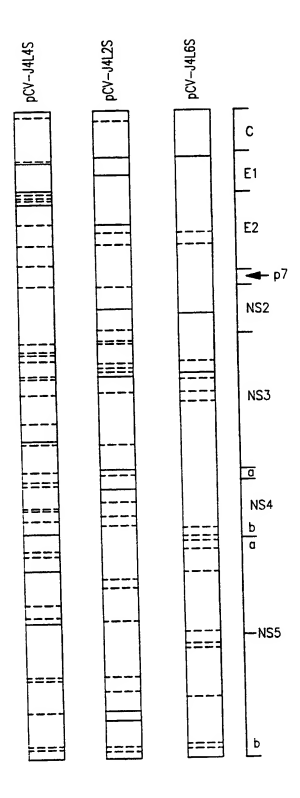


FIG. 12

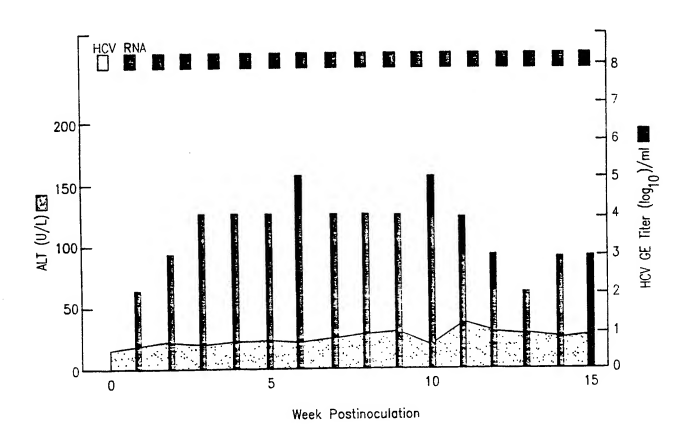


FIG. 13

## HC-J4

				· · · · · · · · · · · · · · · · · · ·	
10	20	30	40	50	
	1234567890				
	TGATGGGGGC				50
	TCTTCACGCA				100
	CCTCCAGGAC				150
	GAGTACACCG				200
	CICAAIGCCI				250
	GIGITGGGIC				300
	GIGCCCCGGG				350
	CICAAAGAAA				400
GGACGICAAG	TICCCGGGGG	GIGGICAGAT	CGTTGGTGGA	GITTACCIGI	450
TGCCGCGCAG	GGGCCCCAGG	TIGGGIGIGC	CCCCCACTAG	GAAGGCITCC	500
GAGCGGICGC	AACCTCGTGG	AAGGCGACAA	CCTATCCCAA	AGGCTCGCCG	550
ACCCGAGGGC	AGGGCCTGGG	CTCAGCCCGG	GTACCCTTGG	CCCCICIAIG	600
GCAATGAGGG	CCTGGGGTGG	GCAGGATGGC	TCCTGTCACC	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	650
CGGCCTAGTT	GGGGCCCCAC	GGACCCCCGG	CGTAGGTCGC	GTAACTIGGG	700
TAAGGICATC	GATACCCTTA	CATGCGGCTT	CCCCGATCTC	ATGGGGTACA	750
TICCGCICGI	CGGCGCCCCC	CTAGGGGGGG	CTGCCAGGGC	CTTGGCACAC	800
CETETCCCGC	TICIGGAGGA	CGGCGTGAAC	TATGCAACAG	GGAACTTGCC	850
COGTTCCTCT	TICICIAICI	TCCTCTTGGC	TCTCCTGTCC	TGTTTGACCA	900
TCCCAGCTTC	CGCTTATGAA	GTGCGCAACG	TGTCCGGGAT	ATACCATGIC	950
ACGAACGACT	GCTCCAACTC	AAGCATIGIG	TATGAGGCAG	CGGACGIGAT	1000
CATGCATACT	CCCGGGIGCG	TGCCCTGTGT	TCAGGAGGGT	AACAGCICCC	1050
GTTGCTGGGT	AGCGCTCACT	CCCACGCTCG	CGGCCAGGAA	TGCCAGCGIC	1100
CCCACTACGA	CAATACGACG	CCACGTCGAC	TTCCTCGTTG	GGACGGCIGC	1150
TTTCTCCTCC	GCTATGTACG	TGGGGGATCT	CIGCGGATCT	ATTTTCCTCG	1200
TCTCCCAGCT	GITCACCITC	TOGCCTCGCC	GGCATGAGAC	AGTGCAGGAC	1250
TGCAACTGCT	CAATCTATCC	CGGCCATGIA	TCAGGICACC	GCATGGCTTG	1300
CGATATGATG	ATGAACIGGI	CACCIACAAC	AGCCCTAGTG	GIGICGCAGI	1350
TECTCOGGAT	CCCACAAGCT	GICGIGGACA	TEGTEGEGGG	GGCCCACTGG	1400
GGAGTCCTGG	CGGGCCTTGC	CIACIATICC	ATGGTAGGGA	ACTOGGCTAA	1450
GGTTCTGATT	GIGGCGCIAC	TCTTTGCCGG	CCTTCACCCC	GAGACCCACA	
CGACGGGGAG	GGIGGCCGGC	CACACCACCI	COGGGIICAC	GICCCITTIC	1550
TCATCTGGGG	CGICICAGAA	AATCCAGCTI	GIGAATACCA	ACGCAGCIG	1600
GCACATCAAC	AGGACTGCCC	TAAATTGCAA	TGACICCCIC	CAAACIGGGI	1650
				COGGIGCCCG	1700
				AGGGGTGGGG	1750
				CCTTATTGCT	1800
				GCAGGIGIGI	
GGTCCAGTGI	ATTGTTTCAC	CCCAAGCCCI	GIIGIGGIGG	GGACCACCGA	1900

FIG. 14A

### HC-J4

10 20	30	40	50	
1234567890 1234567890 123456	_	1234567890	1234567890	
TOGTICOGOT GICCCIACGT ATAGCI				1950
TGCTCCTCAA CAACACGCGT CCGCCA	CAAG (	GCAACIGGIT	CGGCTGTACA	2000
TGCATGAATA GTACTGGGTT CACTAA	GACG '	TGCGGAGGTC	CCCCGIGIAA	2050
CATCGGGGG GICGGIAACC GCACCI	TGAT (	CTGCCCCACG	CACTCCTTCC	2100
GGAAGCACCC CGAGGCTACT TACACA	TAAA	GIGGCIGGGG	GCCCIGGIIG	2150
ACACCIAGGI GCCIAGIAGA CIACOC	ATAC .	AGGCTTTGGC	ACIACCCIG	2200
CACICICAAT TITICCATCT TITAGG				2250
AGCACAGGCT CAATGCCGCA TGCAAT	TGGA	CTCCAGCAGA	GCGCTGTAAC	2300
TIGGAGGACA GGGATAGGIC AGAACI	CAGC	CCCCTCCTCC	TGICIACAAC	2350
AGAGIGECAG ATACIGCCCT GIGCIT	TCAC	CACCCTACCG	GCTTTATCCA	2400
CIGGITICAT CCATCICCAT CAGAAC	ATCG	TGGACGIGCA	ATACCIGIAC	2450
GGIGIAGGT CAGCGTTTGT CTCCT	TIGCA	ATCAAATGGG	AGUACATOCT	2500
GITECTITIC CTICICCIEG CAGACO				2550
TGATGCIGCT GATAGCCCAG GCIGAC				2600
CTCAATGCGG CGTCCGTGGC CGGAGC				2650
GITCITCIGC GCCGCCIGGT ACATTL				2700
CGIATGCITT TTATGGCGIA TGGCC	ECTGC			2750
CCACCACGAG CTTACGCCTT GGACC				2800
1/3/311/211 01/20-0-0-		CTIGICACCA		2850
TGITTCICAC TAGGCICATA TGGIG	FITAC	AATACTITAT		2900
CAGGCGCACA TGCAAGTGTG GGTCC				2950
CGATGCCATC ATCCTCCTCA CGIGT				3000
ACATCACCAA ACTCCTGCTC GCCAT		GCCCGCTCAT		3050
GCTGGCATAA CGAGAGTGCC GTACT				3100
TGCATGCATG TTAGTGCGAA AAGTC				3150
TCTTCATGAA GCTGGGCGCG CTGAC	AGGIA	CGTACGITTA	TAACCAICIT	3200
ACCCCACTGC GGGACTGGGC CCACG	CCCC	CIACGAGACC	TIGOGGIGGO	3250
GGIAGAGCCC GICGICTICT CCGCC	'ATGGA	. GACCAAGGIC	AICACCIGGG	3300
GAGCAGACAC CGCTGCGTGT GGGG	CATCA	TCTIGGGICI	ACCCGICICC	3350
CCCCGAAGGG GGAAGGAGAT ATTT	TGGGA	CCGCIGATA	GICICGAAGG	3400 3450
GCAAGGIGG CGACTCCTIG CGCCC	ATCAC	GGCCIACICC	CAACAAACGC	3 <del>4</del> 50
GGGGGTACT TGGTTGCATC ATCAC	TAGCC	TCACAGGCUC	G GALAALAAL	3550
CACCICCAAC CCCACCIICA ACIC	FITICI	ACCGCAACAC	AAICITICCI	3600
GGCGACCTGC ATCAACGGCG TGTG	CIGGAC	GICTACCA!	r Gererigeri.	
CGAAGACCCT AGCCGGTCCA AAAG	SICCAP	A TCACCCAAA	I. GTATALCANI.	3700
GTAGACCIGG ACCICGICGG CIGG	LAGGCC		- ACCACACAMO	3750
GACACCATGC AGCTGTGGCA GCTO	JGACC.	L LIMCLIGGI	- YZCHANYCH.	3800
CIGATGICAT TCCGGIGCGC CGGC	TYPPE	bbblAblAJA c	אראלו ונאיאים פ	2000

FIG. 14B

### HC-J4

10 20	30	40	50	
1234567890 1234567890			1234567890	
TCCCCAGGC CCGICICCIA	CCTGAAAGGC	TOCTOGGGIG	GICCATIGCT	3850
TIGOCCTICG GGGCACGICG	TEGECETCIT	COGGGGGGGGT	GIGIGCACCC	3900
GGGGGTCGC GAAGGCGGTG	CACTICATAC	COGTTGAGTC	TATGGAAACT	3950
ACCATGOGGT CTCCGGTCTT	CACAGACAAC	TCAACCCCC	CCCCIGIACC	4000
CCACACATTC CAAGIGGCAC	ATCIGCACGC	TCCTACTGGC	AGCGGCAAGA	4050
GCACCAAAGI GCCGGCIGCG	TATGCAGCCC	AAGGGIACAA	GEIGCICGIC	4100
CTGAACCCGT CCGTTGCCGC	CACCTTAGGG	TTTGGGGGGT	ATATGICCAA	4150
GCACACGI ATCGACCCIA	ACATCAGAAC	TGGGGIAAGG	ACCATTACCA	4200
CGGGGGGCIC CATTACGIAC	TOCACCIAIG	GCAAGITCCT	TCCCGACGGT	4250
GCCIGITCIG GGGGGCCTA	TGACATCATA	ATATGIGATG	AGIGCCACIC	4300
AACTGACTCG ACTACCATCT	TEGECATOGG	CACAGICCIG	CACCAAGCGG	<b>435</b> 0
AGACGGCTGG AGCGCGCTC	GICGIGCICG	CCACCGCIAC	ACCICCGGGA	4400
TOGGITACOG TGCCACACCO	CAATATCGAC	GAAATAGGCC	TGTCCAACAA	<b>445</b> 0
TGGAGAGATC CCCTTCTATO	GCAAAGCCAT	CCCCATTGAG	GCCATCAAGG	4500
GGGGAGGCA TCTCATTTTC	TGCCATTCCA	AGAAGAAATG	TCACCACCTC	4550
CCCCAAAGC TGACAGGCC	CGGACTGAAC	GCIGIAGCAT	ATTACCGGGG	4600
CCITGATGIG TCCGTCATAC	CGCCTATCGC	AGACGICGIT	GICGIGGCAA	4650
CAGACGCICT AATGACGGG	TICACCGGCC	ATTTIGACTO	AGTGATCGAC	<b>470</b> 0
TOTALTACAT GTGTCACCC	A GACAGIOGAC	TICAGCIIGG	ATCCCACCIT	4750
CACCATTGAG ACGACGACC	G TGCCCCAAG	i cecedicice	CGCTCGCAAC	4800
CONTACTING AACTIGGCAG	G GGTAGGAGT	GCATCTACAG	GITIGIGACI	4850
MAGGAGAAC GGCCCTCGG	G CATGITCGA	r Terreggiee	TGIGIGAGIG	4900
CTATEACTC GCTGTGCT	r ggratgagc.	r cacgocogci	GAGACCICGG	4950
TTACETTEC GCCTTACCT	A AATACACCA	GGTTGCCCGT	CIGCCAGGAC	5000
CATCTICACT TCTGGGAGA	G CGICTICAC	A GGCCTCACCC	ACATAGAIGC	5050
CACTUCTIC TOCCAGACT	A AACAGGCAG	G AGACAACIII	CCTTACCTGG	5100
TGGCATATCA AGCTACAGT	G TGCGCCAGG	G CICAAGCICO	ACCICCAICG	5150
TOTAL ALATTICAAGI	G TCTCATACG	G CTGAAACCTY	A CACIGCACGG	5200
GYANCACY CIGCIGIAI	A GGCTAGGAG	C CGICCAAAA	r GAGGICATICC	5250
TO CACACO CATA ACTA	A TACATCATG	G CATGCATGI	C GGCIGACCIG	5300
CACCINCTICA CTACCACCI	G GGIGCIGGI	A GCCGGAGIC	TIGCAGCITT	5350
CONTRATAC TOCCTGACO	A CAGGCAGIG	T GGTCATIGIO	G GGCAGGAICA	5400 5450
TOTAL GAAGCCAG	T GROGITOCC	G ACAGGGAAG	r cciciaccag	5 <b>45</b> 0 5500
GAGTICGATG AGATGGAAC	A GIGIGOCIO	'A CAACITOCI	I. ACATOGAGCA	5550 5550
GGGAATGCAG CTCGCCGAC	E AATTCAAGC	A AAAGGCGCT	C GGGLIGIIGC	5550 5600
AAACGGCCAC CAAGCAAGC	OG GAGGCIGCI	G CICCCGIGG	J. GCARTICCAAR	
TGGCGAGCCC TTGAGACC	M CIGGGCGAA	G CACATGIGG	A ATTICATOR	5700
CGGAATACAG TACCTAGC	AG GCTTATCCA	C TOIGCIG	M MALLICUSCUM	2,00

FIG. 14C

### HC-J4

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
TAGCATCATT	GATGGCATTT	ACAGCITCIA	TCACTAGCCC	GCTCACCACC	5750
CAAAACACCC	TCCTGTTTAA	CATCTTGGGG	GGATGGGTGG	CIGCCCAACT	5800
CCCTCCTCCC	AGCGCIGCGI	CAGCITICGI	GGGCGCCGGC	ATCGCCGGAG	5850
CGCCTGTTGG	CAGCATAGGC	CTTGGGAAGG	TGCTCGTGGA	CATCTIGGCG	5900
GGCTATGGGG	CAGGGGTAGC	CGGCGCACTC	GIGGCCITIA	ACCICATCAC	<b>595</b> 0
CGGCGAGGTG	CCCTCCACCG	AGGACCIGGT	CAACITACIC	CCTGCCATCC	6000
TCTCTCCTGG	TECCCIGGIC	GICGGGGICG	TGTGCGCAGC	AATACTGCGT	<b>605</b> 0
CGGCACGTGG	GCCCGGGAGA	GGGGGCTGTG	CAGIGGAIGA	ACCGCCTGAT	6100
ACCGTTCGCT	TOGOGGGGIA	ACCACGICIC	CCCTACGCAC	TATGIGCCIG	6150
AGAGCGACGC	TGCAGCACGI	GICACICAGA	TCCTCTCTAG	CCTTACCATC	6200
ACTCAACTGC	TGAAGOGGCT	CCACCAGIGG	ATTAATGAGG	ACIGCICIAC	6250
GCCATGCTCC	GCTCGTGGC	TAAGGGATGT	TTGGGATTGG	ATATGCACGG	6300
TGITGACIGA	CITCAAGACC	TGGCTCCAGT	CCAAACTCCT	GCCGCGGTTA	6350
CCGGGAGTCC	CITICCIGIC	ATGCCAACGC	GGGTACAAGG	CACICICCC	6400
GGGGGALGGC	ATCATGCAAA	CCACCIGCCC	ATGCGGAGCA	CAGATOGCCG	6450
GACATGICAA	AAACGGTTCC	ATGAGGATCG	TAGGGCCTAG	AACCIGCAGC	6500
AACACGIGG	ACGGAACGIT	CCCCATCAAC	GCATACACCA	CCCCACCTIC	6550
CACACCCICC	COGGGGGCCA	ACTATICCAG	GGCGCTATGG	CEGGLIGGCLC	<b>660</b> 0
CIGAGGAGIA	CGIGGAGGII	ACCCGIGICG	GGGATTTCCA	CIACGIGACG	6650
GGCATGACCA	A CIGACAACGI	' AAAGTGCCCA	TGCCAGGIIC	CGGCCCCCGA	<b>670</b> 0
ATTCTTCACC	GAGGIGGATC	GAGTGCGGTT	GCACAGGIAC	GCICCGGGGI.	6750
GCAAACCICI		GACGICACGI			6800
TACTIGGIC				ACGIAACAGI	6850
GCTTACTTC	ATGCTCACCG				6900
AGCGTAGGC.				CICATCAGCT	6950
AGCCAGTIG.				CCCACCATGA	7000
				CGGCAGGAGA	7050
	A CATCACICGO				7100
				AGATATOOGT	7150
				GCGTTGCCCA	7200
				CIGGAAGGAC	
				CACCTACCAA	7300
				GICCIGACAG	7350
				A GACCTICGGI	7400
				CCCTTCCTGA	
				TOGIACICCI	7500
				r CAGCGACGGG	7550
TCTTGGTCT	'A CCGTGAGTE	A GGAGGCTAG	r gaggaigic	G TETGETGETE	7600

FIG. 14D SUBSTITUTE SHEET (RULE 26)

## HC-J4

10	20	30	40	50	
1024567990	1234567890				
123430703U	ACGIGGACAG	GGCCTGAT	CACGCCATGC	GCTGCGGAGG	7650
	GCCCATCAAC				7700
	ACGCCACAAC				7750
	GACAGATTGC				7800
	CAACCCCAAC				7850
	CCTGCAAGCT				7900
	GCAAAGGACG				7950
ACAMOCCIALOGG	CGIGIGGGAG	CACTUGCIGG	AAGACACIGA	AACACCAATT	8000
CACACCACTA	TCATGGCAAA	AAGTGAGGIT	TICIGOGICC	AACCAGAGAA	8050
CHILATORICA	AAGCCAGCIC	COTTATOGT	ATTCCCAGAC	CIGGGAGIIC	8100
	GAAGATGGCC				8150
CIGIAICCE	GCICCICATA	CGATTICAA	TACTOCCCCA	AGCAGCGGGT	8200
Column	GIGAATACCT	GGAAATCAAA	GAAATGCCCT	ATGGGCTTCT	8250
CAMATICCIO	CCCCIGITIT	GACTCAACGG	TCACTGAGAG	TGACATTCGT	8300
CATATORNET	CAATTIACCA	ATGITGICAC	TTGGCCCCCG	AGGCCAGACA	8350
GITGENERAL	TOGOTCACAG	AGCGGCTTTA	CATCGGGGGT	CCCCTGACTA	8400
ACTUADADAG	GCAGAACTGC	GGITATCGCC	GGTGCCGCGC	AAGTGGCGTG	8450
CTCACCACTA	GCTGCGGTAA	TACCCICACA	TGTTACTTGA	AGGCCACTGC	8500
	GCTGCAAAGC				8550
ACCOUNTS!	CGITATCIGI	GAAAGOGCGG	GAACCCAGGA	GGATGCGGCG	8600
COTACAC	CCTTCACGGA	GGCTATGACT	AGGIATICCG	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	8650
GATOGOO	CAACCAGAAT	ACGACCIGGA	GCTGATAACA	TCATGTTCCT	<b>870</b> 0
CAATGIGIC	AGTOGOGCAC	GATGCATCIG	GCAAAAGGGT	ATACTACCIC	8750
ACCCTGACC	CACCACCCC	CCTTGCACGG	GCTGCGTGGG	AGACAGCTAG	8800
ACACACTOCA	ATCAACICIT	GGCTAGGCAA	TATCATCATE	TATGCGCCCA	8850 ~
CCCTATGGG	AAGGATGATT	CIGATGACIO	ACTITICIO	CATCCTICIA	8900
GCTCAAGAG	AACTIGAAAA	AGCCCTGGAT	TGICAGAICI	ACGGGGCTTG	8950
CTACTCCATT	GAGCCACTIG	ACCIACCICA	GATCATIGAA	CGACTCCATG	9000
GICTIAGCG	ATTTACACTO	CACAGTTACT	CICCAGGIGA	CATCAATAGG	9 <b>05</b> 0
GIGGCTICA	CCCTCAGGAA	ACTIGGGGIA	CCACCCTIGO	GAACCIGGAG	9100
ACATOGGGC	AGAAGIGICC	GCGCTAAGCI	ACIGICCCAC	GGGGGGAGGG	9150
CCGCCACTI	TGGCAGATAC	CICITIAACI	C GGGCAGIAAC	GACCAAGCTT	9200
AAACTCACT	CAATCCCGG	CGCGTCCCAL	G CIGGACTIG	CIGGCIGGIT	9250
CGTCGCTGG	TACAGOGGG	GAGACATATA	A TCACAGOCTO	TCICGIGCCC	9300
GACCCCCCT	GITICOGITO	TOCCIACIO	TACITICIG	r aggggiaggc	9350
ATTTACCTG	TOCCCAACCO	ATGAACGGG	G AGCTAACCA	TOCAGGCCIT	9400
AAGCCATTT	CIGITITITE	TTTTTTTT	r TTTTTTTTT	TCTTTTTTT	9450
TTTCTTTCC	T TREETTERT	TITIOCITI	TITTICCI.	CITTAATGGT	9500

FIG. 14E

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
GGCTCCATCT	TAGCCCTAGT	CACGGCTAGC	TGTGAAAGGT	CCGIGAGCCG	9550
CATGACTGCA	GAGAGIGCIG	ATACIGGCCT	CICIGCAGAT	CAIGI	9595

FIG. 14F

10	20	30	40	50	
1234567890 12345	67890	1234567890	1234567890	1234567890	
MSTNIPKPORK TKRNI	NRRPO	DVKFPGGGQI	VGGVYLLPRR	GPRLGVRAIR	50
KASERSOPRG RROPI	PKARR	PEGRAWAQPG	YPWPLYGNEG	LGWAGWLLSP	100
RESRPSWEPT DPRRE	SRNLG	KVIDILICGF	ADIMGYIPLV	<b>GAPLGGAARA</b>	150
TAHEVRVLED GVNYA	TGVLP	<b>GCSFSIFLLA</b>	LLSCLTIPAS	AYEVRNVSGI	200
VHITNICSNS SIVYE	ZADVI	MHIPGCVPCV	<b>QEGNSSROW</b>	ALIPILAARN	250
ASTETTTIRE HVDLI	MGIAA	FCSAMYVGDL	CGSIFLVSQL	FIFSPRRHEI	300
VODONCSTYP GHV90	HRMAW	<b>IMMNWSPIT</b>	ALVVSQLLRI	POAVVIMVAG	350
AHMENTAGIA YYSM	CIWAK	VLIVALLFAG	VDGEIHITGR	VACHITSGFT	400
ST.FSSGASOK IOLM	VIIVGSW	HINRIALNON	DSLQIGFFAA	LFYAHKFNSS	450
GOPERMASCR PIDWE	FACGNG	PITYTKPNSS	DORPYCWHYA	PRPCGVVPAS	500
OVICEPVYCET PSPV	WGIID	RSGVPTYSWG	ENEIDVMLIN	NIRPPQGWF	550
CCIWMNSIGF TRICO	<b>GPPCN</b>	IGGVENRILI	CPIDCFRKHP	EATYTKOGSG	600
PWI TPRCLVD YPYRI	WHYPC	TLNFSIFKVR	MYVGGVEHRL	NAACIWIRGE	650
RONLEDRORS ELSPI	LISTT	EWOILPCAFT	TLPALSIGLI	HLHQNIVDVQ	700
VINCUCSARY SEAT	<b>WEYIL</b>	LIFILLADAR	VCACLIMMIL	IAQAEAALEN	750
TIMINAASVA GAHG	ILSFLV	FFCAAWYIKG	RLAPGAAYAF	YGWPLLLLL	800
TAT.PPRAYAT, DREM	AASCGG	AVLVGLVFLT	LSPYYKVFLT	RLIWWLQYFI	850
TRAFAHMOW VPPL	WROGR	DAIILLICAV	HPELIFDITK	LLLAILGPLM	900
VI CACTURVE YEVR	AOGLIR	ACMLVRKVAG	CHYVQMVFMK	LGALIGIYVY	950
NHI TPI RIWA HAGL	RDLAVA	VEPVVFSAME	TKVITWGADI	AACGDIIIGL	1000
PVSARRCKEI FLGP	ADSLEG	QGWRLLAPIT	AYSQQIRGVL	GCIITSLIGR	1050
DKNOVEGEVO VVSI	ATQSFL	ATCINGVCWI	VYHGAGSKIL	AGPKGPIIQM	1100
YTNIDIDIIG WOAP	PGARSM	TPCSCGSSDL	, YLVIRHADVI	PVRRRGDSRG	1150
STISPRPVSY LKGS	SGGPLL	CPSGHVVGVF	'RAAVCIRGVA	KAVDFIPVES	1200
METIMRSPVF TINS	TPPAVE	OTFQVAHLHA	, PIGSGKSTKV	PAAYAAQGYK	1250
VLVLNPSVAA TLGF	GAYMSK	AHGIDPNIRI	GVRITITIGGS	TTYSTYCKFL	1300
ADGGCSGGAY DIII	CDECHS	TDSTTILGIC	TVLDQAETAC	ARLVVLATAT	1350
PPGSVIVPHP NIE	IGLSW	GEIPFYGKAI	PIEAIKGGR	LIFCHSKKKC	1400
DELAAKLIGL GLNA	VAYYRC	LDVSVIPPIC	DVVVVATDAI	MIGFIGDEDS	1450
VIDONICVIQ TVDE	SLDPIE	TIETTIVPQI	) AVSRSQRRGF	R TGRGRSGIYR	1500
FATPCERPSG MEDS	SVLCE	YDAGCAWYEI	_ TPAETSVRLF	R AYLNIPGLIPV	1550
CODHLEFWES VFTC		A HFLSQIKQAC	DIFPYLVAY(	ATVCARAQAP	1600
PPSINTYMINKC I.TRI	KPILK	PIPLLYRLG	A AĞMEATTIHI	PITKYIMACMS	7,020
ADLEVVISIW VLV	3GVLAAI	_ AAYCLITGS	/ VIVGRIILS	S KPAVVPLKEV	1700 1750
LYQEFDEMEE CAS	DLAAID	O GMOLAEOFK	XALGLIQIA'	L KÜAFAAAFAA	
ESKWRALETF WAK	HMMTI:	S GIQYLAGLS	r LPGNPAIASI	- CICL CANAN	
LTTQNILLFN ILG	QAAVWE	L APPSAASAF	V GAGLAGAAV	2 DICHIMANAN	1900
ILAGYGAGVA GAL	VAFKVM	S GEVPSTEDL	V NILPAILSP	ALVVOVVLAA 6	1500

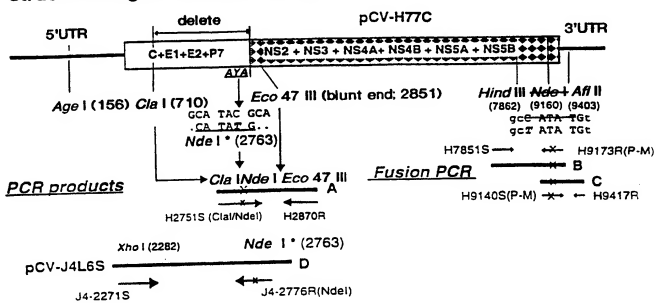
FIG. 14G

SUBSTITUTE SHEET (RULE 26)

10	20	30	40	50	
1234567890 12	234567890	1234567890	1234567890	1234567890	
TI RRHVGPGE G	AVOWMNRLI	<b>AFASRGNHVS</b>	PIHYVPESDA	AARVIQILSS	1950
LTTTOLLKRL H	WINEDCST	PCSGSWLRDV	MDWICIVLID	FKIWLQSKLL	2000
PRLPGVPFLS CO	DRGYKGVWR	GDGIMQITCP	CGAQIAGHVK	NGSMRIVGPR	2050
TOSNIWHGIF P	NAYTIGPC	TPSPAPNYSR	ALWRVAAEEY	VEVIRVGDFH	2100
VATIONITIAN KO	CPCOVPAPE	FFTEVDGVRL	HRYAPACKPL	LREDVIFQVG	2150
INDYINGSOL PO	CEPEPDVIV	LISMLIDPSH	TTAETAKRRL	ARGSPPSLAS	2200
SSASOLSAPS L	KATCITHHD	SPDADLIEAN	LLWROEMGGN	TIRVESENKV	2250
VII DSFEPLH A	PEDERELSV	AAEILRKSRK	<b>FPSALPIWAR</b>	PDYNPPLLES	2300
WKDPDYVPPV V	HCCPLPPIK	APPIPPPRRK	RIVVLIESW	SSALAELATK	2350
TEGSSGSSAV D	SCIPITALPD	LASDDGDKGS	DVESYSSMPP	LEGEPGOPDL	2400
SDGSWSTVSE E	ASEDVVCCS	MSYTWICALI	TPCAAEESKL	PINPLSNSLL	2450
RHHNMVYATT S	RSASLROKK	VIFDRLQVLD	DHYRDVLKEM	KAKASIVKAK	2500
TISTEFACKI T	PPHSAKSKF	GYGAKDVRNL	SSRAVNHIRS	WEDLLEDIE	2550
TPTTTTTMAK S	EVECVOPEK	GGRKPARLIV	FPDLGVRVCE	KMALYDVVST	2 <b>60</b> 0
T.POAVMGSSY G	FOYSPKORV	EFLVNIWKSK	KCPMGFSYDI	RCFDSIVIES	2650
DIRVEESIYO C	CDLAPEARQ	AIRSLIERLY	IGGPLINSKG	QNCGYRRCRA	<b>270</b> 0
SGVLTTSCGN I	LICYLKATA	ACRAAKLQDO	MINGDOLV	VICESAGIQE	2750
DADATRAFTE A	MIRYSAPPG	DPPOPEYDLE	LITSCSSNVS	VAHDASGKRV	2800
YYITRDPITP I	ARAAWETAR	HIPINSWLGN	IIMYAPILWA	RMILMIHFFS	2850
TTI AOFOLEK A	LDCOTYGAC	YSIEPLDLPC	) ITERLHGLSA	FILHSYSPGE	2900
TARKASATEK I	GIPPLRIWE	HRARSVRAKI	_ LSQGGRAAT(	GRYLFNWAVR	<b>295</b> 0
TKLKLTPIPA A	ASOLDLSGWE	VAGYSGEDIY	HSLSRARPR	FPLCILLISV	<b>300</b> 0
GVGIYLLPNR					<b>301</b> 0

**FIG. 14H** 

#2. Strategy for constructing chimeric clone of HCV (pH77CV-J4) which contains the nonstructural region of strain H77 and the structural region of strain HC-J4



- 1. Fragment A, B, C and D; PCR amplification from pCV-H77C or pCV-J4L6S
  - Fragment A; additional Cla I site, artificial Nde I site induced by a single mutation (C→T at nt 2765 of H77C) and authentic Eco47 III site
  - Fragment B and C; eliminated Nde I site by a single mutation within the primers (C→T at nt 9158 of H77C), and fusion PCR with both fragments
  - Fragment D; artificial Nde I site induced by 2 point mutations within the primer (T→A at nt 2762 and C→T at nt 2765 of J4L6S)
- 2. TA cloning of PCR products
- 3. Sequence analysis
- 4. Cloning of Fragment A (Cla I-Eco 47III ) and Fragment B/C (Hind III-Afl II ) with correct sequence into pCV-H77C
- 5. Complete sequence analysis of new cassette vector [pH77CV], into which the structural regions of different genotypes can be inserted.
- 6. Cloning of Fragment-Age I/Xho I (cut out from pCV-J4L6S) and Fragment D (Xho I-Nde I) with correct sequence into the new cassette vector; 3 piece ligation
- 7. Complete sequence analysis of 1a+1b chimera [pH77CV-J4]
- 8. In vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee

FIG. 15

GCCAGCCCC TGATGGGGCC GACACTCCAC CATGAATCAC TCCCCTGTGA	50
GGAACTACTG TCTTCACGCA GAAAGCGTCT AGCCATGGCG TTAGTATGAG	100
TGTCGTCCAG CCTCCAGGAC CCCCCCTCCC GGGAGAGCCA TAGTGGTCTG	150
CCGAACCCCT CACIACACCG GAATTCCCAG GACCACCCCG TCCTTTCTTG	200
CATCAACCCG CTCAATGCCT GGAGATTIGG GCGIGCCCCC GCGAGACIGC	250
TAGCOCAGIA GIGITGGGIC GOCAAAGGOC TIGIGGIACT GCCIGATAGG	300
GIGCIIGOGA GIGCCCCGGG AGGICIOGIA GACOGIGCAC CATGAGCACG	350
AATCCIAAAC CTCAAAGAAA AACCAAACGT AACACCAACC GCCGCCCACA	400
GCACGICAAG TICCCGGGGG GIGGICAGAT CGTIGGIGGA GITIACCIGI	450
TECCECECAG GEGCCCCAGG TIGGGIGIGC GCGCGACIAG GAAGGCTICC	500
GAGOGGICGC AACCICGIGG AAGGOGACAA CCIATCCCAA AGGCICGCCG	550
ACCOGAGESC AGGEOCIGES CICAGOCOGG GIACOCTIGG COCCICIATG	600
GCAATGAGGG CCTGGGGTGG GCAGGATGGC TCCTGTCACC CCGGGGCTCC	650
CCCCCTAGTT CCCCCCCAC CCACCCCCCG CGTAGGTCGC GTAACTTGGG	700
TAAGGICATC GATACCCTTA CATGCGGCTT CGCCGATCTC ATGGGGTACA	750
TICCGCICGI CGGCGCCCC CTAGGGGGG CTGCCAGGGC CTTGGCACAC	800
GGIGICCGGG TICIGGAGGA CGGCGIGAAC TAIGCAACAG GGAACIIGCC	850
COGFIGCICI TICICIATOT TOCICTICOC TOTOCCACCA	900
TCCCAGCITC CGCITATGAA GIGCGCAACG TGTCCGGGAT ATACCATGTC	950
ACCAACCACT CCTCCAACTC AAGCATTGTG TATGAGGCAG CGGACGTGAT	1000
CATGCATACT CCCGGGIGCG TGCCCTGTGT TCAGGAGGGT AACAGCTCCC	1050
GITCCICCOT ACCCCICACT CCCACCCICG CCCCACCAA TCCCACCGIC	1100
CCCACTACGA CAATACGACG CCACGTCGAC TICCTCGTTG GGACGCTGC	1150
TITCICCICC GCIAIGIACG TGGGGGAICT CIGCGGAICT ATTITCCICG	1200
TCTCCCAGCT GITCACCTTC TCGCCTCGCC GGCATGAGAC AGTGCAGGAC	1250
TGCAACIGCT CAATCIATCC CGGCCATGIA TCAGGICACC GCATGGCITG	1300
GGATATGATG ATGAACTGGT CACCTACAAC AGCCCTAGTG GTGTCGCAGT	1350
TECTICOGEAT COCACAAGCT GTOGTGGACA TEGTGGCGGG GGCCCACTGG	1400
GGAGICCIGG CGGCCFIGC CIACIATICC AIGGIAGGA ACIGGCCIAA	1450
GGITCIGATT GIGGCGCIAC TCTTTGCCGG CGTTGACGGG GAGACCCACA	1500
CGACCGGGAG GGIGGCCGGC CACACCACCT CCGGGITCAC GICCCITTIC	1550
TCATCIGGG CGICICAGAA AATCCAGCTT GIGAATACCA ACGCAGCIG	
GCACATCAAC AGGACTGCCC TAAATTGCAA TGACTCCCTC CAAACTGGGT	1650
TCTTTGCCGC GCIGTTTTAC GCACACAAGT TCAACTCGTC CGGGTGCCCG	1700
CACCCATICS CCACCICCCG CCCCATICAC TGGITCGCCC AGGGIGGGG	1750
CCCCATCACC TATACTAAGC CTAACAGCTC GGATCAGAGG CCTTATTGCT	1800

FIG. 16A

GECATTACEC GOCTOGACCG TGTGGTGTCG TACCOGCGTC GCAGGTGTGT	1850
GGICCAGIGT ATTGITTCAC CCCAAGCCCT GITGIGGIGG GGACCACCGA	1900
TOGITOCGGT GICCCTACGT ATACCTGGGG GCAGAATGAG ACAGACGTGA	1950
TGCTCCTCAA CAACACGCGT CCGCCACAAG GCAACTGGTT CGGCTGTACA	2000
TOCATCAATA GIACTOCCIT CACTAACACG TOCGCACGIC CCCCGIGIAA	2050
CATCEGEGE GIOSTAACC GCACCITGAT CIGCCCCACG GACTECITCC	2100
GGAAGCACCC CGAGGCTACT TACACAAAAT GIGGCTCGGG GCCCIGGITG	2150
ACACCIAGGI GCCIAGIAGA CIACCCATAC AGGCTTIGGC ACIACCCCIG	2200
CACICICAAT TITICCATCI TIAAGGITAG GAIGIATGIG GGGGGGGGG	2250
AGCACAGGCT CAATGCCGCA TGCAATIGGA CTCGAGGAGA GCGCTGTAAC	2300
TIGGAGGACA GGGATAGGIC AGAACICAGC CCGCIGCIGC TGICIACAAC	2350
AGAGIGGCAG ATACIGCCCT GIGCITICAC CACCCTACCG GCTTIATCCA	2400
CIGGITICAT CCATCICCAT CAGAACATCG TGGACGIGCA ATACCIGIAC	2450
GGIGIAGGGT CAGCGTTIGT CICCTTIGCA ATCAAATGGG AGIACATCCT	2500
GITGCITTIC CTICICCIGG CAGACGCGCG CGIGIGIGCCC TECTTGIGGA	2550
TGATGCIGCT GATAGCCCAG GCIGAGGCCG CCTTAGAGAA CTTGGIGGIC	2600
CTCAATGCGG CGTCCGTGGC CGGAGCGCAT GGTATTCTCT CCTTTCTTGT	2650
GITCTTCTGC GCCGCCTGGT ACATTAAGGG CAGGCTGGCT CCTGGGGGGG	2700
CGIAIGCITT TIAIGGCGIA TGGCCGCIGC TCCIGCICCT ACIGGCGTIA	2750
CCACCACGAG CATATGCACT GGACACGGAG GTGGCCGCGT CGTGTGGCGG	2800
CGFIGITCIT GICGGGTIAA TGGCGCIGAC TCIGICGCCA TATTACAAGC	2850
CCTATATCAG CIGGICCATG TGGIGGCTIC AGPATITICT GACCAGAGIA	2900
GAAGCGCAAC TGCACGTGTG GGTTCCCCCC CTCAACGTCC GGGGGGGCG	2950
CGATGCCGIC ATCTIACICA TGIGIGIAGI ACACCCGACC CIGGIATITG	3000
ACATCACCAA ACTACTCCTG GCCATCTTCG GACCCCTTTG GATTCTTCAA	3050
COCAGTTICC TTANAGICCC CTACTTCGIG CCCGTTCAAG CCCTTCTCCG	3100
CATCIGOGOG CIAGOGOGGA AGATAGOCGG AGGICATTAC GIGCAAATGG	3150
CCATCATCAA GTTAGGGGG CTTACTGGCA CCTATGTGTA TAACCATCTC	3200
ACCCCICTIC GAGACIGGC GCACAACGC CIGCGAGAIC IGCCCGIGGC	3250
TGIGGAACCA GICGICTICT CCCGAATGGA GACCAAGCIC ATCACGIGGG	3300
GGGCAGATAC CGCCGCGIGC GGIGACATCA TCAACGGCIT GCCCGICICT	3350
GCCCGIAGGG GCCAGGAGAT ACIGCITGGG CCAGCCGACG GAATGGICIC	3400
CAAGGGGG ACGITGCTGG CGCCCATCAC GGCGTACGCC CAGCAGACGA	3450
CACCCTCCT ACCCIGIATA ATCACCACCC TCACTGCCCG GCACAAAAAC	3500
CAAGIGGAGG GIGAGGICCA GAICGIGICA ACIGCIACCC AAACCIICCI	3550
GCCAACGIGC ATCAATGGGG TATGCIGGAC TGICTACCAC GGGGCCGGAA	3600

FIG. 16B

CGAGGACCAT CGCATCACCC AAGGGICCIG TCATCCAGAT GIATACCAAT	3650
GIGGACCAAG ACCITGIGGG CIGGCCCGCT CCICAAGGIT CCCGCICATT	3700
GACACCCIGI ACCIGCECT CCICGGACCT TIACCIGGIC ACGAGGCACG	3750
CCGATGICAT TCCCGIGCGC CGGCGAGGIG ATAGCAGGGG TAGCCIGCTT	3800
TOGOCCOGGO CCATTICCIA CITGAAAGGO TOCTOGGGG GICCGCIGIT	3850
GIGCCCCCC GGACACCCC TGGCCCIATT CAGGCCCGC GIGICCACCC	3900
GIGGAGIGGC TAAAGCCGIG GACTTTATCC CIGIGGAGAA CCIAGGGACA	3950
ACCATGAGAT CCCCGGIGIT CACGGACAAC TCCTCTCCAC CAGCAGIGCC	4000
CCAGAGCTTC CAGGTGGCCC ACCTGCATGC TCCCACCGGC AGCGGTAAGA	4050
GCACCAAGGT CCCGGCTGCG TACGCAGCCC AGGGCTACAA GGTGTTGGTG	4100
CTCAACCCCT CIGITGCIGC AACGCIGGGC TTTGGIGCIT ACATGICCAA	4150
GCCCATGGG GTTGATCCTA ATATCAGGAC CGGGGTGAGA ACAATTACCA	4200
CTEGCAGOOC CATCACGIAC TOCACCIACG GCAAGITCCT TGCCGACGCC	4250
GETTECTICAG CAGGICCITA TCACATAATA ATTIGICACG AGICCCACIC	4300
CACCEATECE ACATECATET TEGGCATEGG CACTETECTT GACCAAGCAG	4350
ACACTECTES CECCAGACIG GIIGICCICG CCACTGCTAC CCCICCGGC	4400
TOTALICATIC TETECCATCE TAACATOGAG GAGGITGCIC TETECACCAC	4450
COCACACATO COCHTHACG GCAAGGCHAT COCCCIOGAG GIGATCAAGG	4500
CECCAACACA TCTCATCTTC TGCCACTCAA AGAAGAAGTG CGACGAGCTC	4550
COCCUPACE TEGICGEATT GESCATCAAT GEOGIGGEET ACTACEGEG	4600
TOTTELOGIG TOTGICATOO OGACCAGOGG CGATGITGIC GIOGIGICGA	4650
COCATECTOT CATGACTEGO TITACOGGOG ACTTOGACTO TETGATAGAC	4700
TECAACACET GIGICACTCA GACAGICGAT TICAGCCTIG ACCCTACCTT	4750
TACCATICAG ACAACCACGC TCCCCCAGGA TGCTGTCTCC AGGACTCAAC	4800
GCCGGGGCAG GACTGGCAGG GCGAAGCCAG GCATCTATAG ATTTGTGGCA	4850
CONTRACT COCCUCOGG CATGITOGAC TOGICOGICC TCIGIGAGIG	4900
CTATEACTO COCIGIGATE GENERAL CACGOCOCC GAGACIACAG	4950
TURGECURCE ACCETACATE AACACCCCC GCCTTCCCGT GTGCCAGGAC	5000
CARCITICA ATT TITUTACEAGG CGICITTEAGG CGCCICACIC ATATAGATGC	5050
CACHITITIA TOTCACAA ACCACAGIGG GGAGAACITT OCTTACCIGG	5100
TRACCETACIA AGCCACCGIG TGCGCTAGGG CTCAAGCCCC TCCCCATCG	5150
TOTALOGICA TETEGRAGIG TITGATOCCO CTPARACOCA COCIOCATEG	5200
COCALCACOC CTCCTATACA GACTGCCCCC TGTTCACAAT GAAGTCACCC	5250
TEACCEACCE AATCACCAAA TACATCATGA CATGCATGIC GCCCACCIG	5300
CACCINCINA CGACCACCIG GGIGCICGIT GGCGGGGICC IGGCIGCICI	5350
GCCCCCGTAT TGCCTGTCAA CAGGCTGCGT GGTCATAGTG GGCAGGATCG	5400
<del></del>	

FIG. 16C

TCTTGTCCGG GAAGCCGGCA ATTATACCTG ACAGGGAGGT TCTCTACCAG	5450
TCTIGICOG GARCOGCA ATTATACCIO ACABOGASSI ICIONACCA	5500
CAGTICGAIG ACATGGAAGA GIGCICICAG CACTIACOGT ACATCGAGCA	5550
AGGGATGATG CTCGCTGAGC AGTTCAAGCA GAAGGCCCTC GGCCTCCTGC	5600
AGACCGCGTC CCGCCATGCA GAGGITATCA CCCCTGCTGT CCAGACCAAC	5650
TGGCAGAAAC TCGAGGICIT TIGGGCGAAG CACAIGIGGA ATTICATCAG	
TGGGATACAA TACTTGGGGG GCCIGTCAAC GCTGCCTGGT AACCCCGCCA	5700 5750
TIGCTICATT CATGCCTTTT ACAGCIGCCG TCACCAGCCC ACTAACCACT	5750
GGCCAAACCC TCCTCTTCAA CATATTGGGG GGGTGGGTGG CTGCCCAGCT	5800
CCCCCCCC GGICCCCIA CIGCCITIGI GGGIGCIGGC CIACCIGGCG	5850
CCGCCATOGG CAGCGITGGA CIGGGGAAGG TCCTCGIGGA CATTCTIGCA	5900
GGGIATGGCG CGGGCGTGGC GGGAGCTCTT GTAGCATTCA AGATCATGAG	5950
COGTIGAGGIC CCCTCCACGG AGGACCTGGT CAATCTGCTG CCCGCCATCC	6000
TCICGCCIGG ACCCCTTGTA GICGGIGIGG TCIGCGCAGC AATACIGCGC	6050
COCCACGITG CCCCGCCCA CCCCCCAGIG CAATGGATGA ACCCCCTAAT	6100
ACCUTICACO TOCOGAGA ACCATATTO COCCACACAC TACATACA	6150
AGAGCGATGC AGCCGCCCC GTCACTGCCA TACTCAGCAG CCTCACTGTA	6200
ACCCAGCICC TGAGGCGACT GCATCAGIGG ATAAGCICGG AGIGIACCAC	6250
TOCATGCICC GGITCCIGGC TAAGGGACAT CIGGGACIGG ATATGCGAGG	6300
TOTTCACCEA CITTAAGACC TGGCIGAAAG CCAAGCICAT GCCACAACIG	6350
COURTEATUR CONTIGUES CIGOCARCOR GEGIATARES GEGICIERCE	6400
AGGAGACGC ATTATGCACA CICGCIGCCA CIGIGGAGCT GAGATCACIG	6450
CACATETCAA AAACGEGACG ATGAGGATCG TCGGTCCTAG GACCTGCAGG	6500
AACATGIGGA GIGGGACGIT CCCCATIAAC GCCIACACCA CGGCCCCIG	6550
TACTOCCCTT CCTGCGCCGA ACTATAAGTT CGCGCTGTGG AGGGTGTCTG	6600
CAGAGGAATA OGIGGAGATA AGGCGGGIGG GGGACTICCA CIACGIAICG	6650
CCTATGACTA CIGACAATCT TAAAIGCCCG TGCCAGATCC CATCGCCCGA	6700
ATTITUTCACA GAATIGGACG GGGIGCGCCT ACACAGGITT GCGCCCCCTT	6750
CLACOCTT CCTCCCCAG GACGTATCAT TCAGAGTAGG ACTCCACGAG	6800
TACCOCATICS GETCGCAATT ACCTIGCGAG CCCGAACCGG ACGTAGCCGT	6850
CHICACETIC ATTECTCACIG ATCCCICCCA TATAACAGCA GAGGGGGCCG	6900
GEAGAAGUT GECEAGAGG TCACCCCTT CIATGGCCAG CICCICGGCT	6950
ACCIACTET COCCICCATO TOTCAAGGCA ACTIGCACOG CCAACCAIGA	7000
CHOCOTICAL GOOGAGCICA TAGAGGCIAA CCICCIGIGG AGGCAGGACA	7050
TEXTOTAL CATCACCAGG GITGAGICAG AGAACAAAGT GGIGATICIG	7100
CACTOCTUCE ATCCCCTTGT CGCAGAGGAG GATGAGCGGG AGGICICUGI	7150
ACCIGCAGAA ATTCIGCGGA AGICICGGAG ATTCGCCCGG GCCCIGCCCG	7200

FIG. 16D

TCTGGGCGCG GCCGGACTAC AACCCCCCGC TAGTAGAGAC GTGGAAAAAG	7250
CCTGACTACG AACCACCTGT GGTCCATGGC TGCCCGCTAC CACCTCCACG	7300
GICCCCICCT GIGCCICCGC CICGGAAAAA GCGIACGGIG GICCICACCG	7350
AATCAACCCT ATCTACTGCC TTGGCCGAGC TTGCCCACCAA AAGTTTTGGC	7400
AGCICCICAA CITCOGGCAT TACGGGGGAC AATACGACAA CATCCICIGA	7450
GCCCGCCCT TCTGGCTGCC CCCCGACTC CGACGITGAG TCCTATTCTT	7500
CCATGOCCC CCTGGAGGG GAGCCTGGGG ATCCGGATCT CAGCGAGGG	7550
TCATGGICGA CGGICAGIAG TGGGGCCCAC ACGGAAGAIG TCGIGIGCIG	7600
CICAATGICT TATTCCIGGA CAGGCGCACT CGICACCCCG TGCGCIGGGG	7650
AAGAACAAAA ACTGCCCATC AACGCACTGA GCAACTCGTT GCTACGCCAT	7700
CACAATCIGG TGIATICCAC CACTICACGC AGIGCTIGCC AAAGGCAGAA	7750
GAAAGICACA TITGACAGAC TGCAAGITCT GGACAGCCAT TACCAGGACG	7800
TGCTCAAGGA GGTCAAAGCA GCGCCGTCAA AAGTGAAGGC TAACTTGCTA	7850
TCCGIAGAGG AAGCITGCAG CCIGACGCCC CCACATTCAG CCAAATCCAA	7900
GITTGGCTAT GGGGCAAAAG ACGTCCGTTG CCATGCCAGA AAGGCCGTAG	7950
CCCACATCAA CICCGIGIGG AAAGACCTIC TOGAAGACAG TGIAACACCA	8000
ATAGACACTA CCATCATGGC CAAGAACGAG GITTICIGCG TICAGCCIGA	8050
CAACCCCCT CGIAACCCAG CICGICICAT CGIGITCCCC CACCTCCCCC	8100
TECECCIGIG CGAGAGATG CCCCIGIACG ACGIGGITAG CAACCICCCC	8150
CIGGOCGIGA TEGGAAGCIC CIACGGATIC CAATACICAC CAGGACAGCG	8200
GETTGAATTC CTCGTGCAAG CGTGGAAGTC CAAGAAGACC CCGATGGGT	8250
TCTCGIATGA TACCCGCTGT TTTGACTCCA CAGTCACTGA GAGCGACATC	8300
CGTACGGACG AGGCAATTTA CCAATGTTGT GACCTGGACC CCCAAGCCCG	8350
CGIGGCCATC AAGTCCCTCA CIGAGAGGCT TIAIGITGGG GGCCCTCTTA	8400
CCAATTCAAG GGGGAAAAC TGCGGCTACC GCAGGTGCCG CGCGAGCGC	8450
GTACIGACAA CTAGCIGIGG TAACACCCIC ACTIGCIACA TCAAGGCCCG	8500
GECAGCCIGT CGAGCCGCAG GGCTCCAGGA CTGCACCATG CTCGTGTGTG	8550
CCCACCACTT AGICGITATC TGICAAAGIG CGGGGGICCA GCAGGACGCG	8600
GCCAGCCIGA CAGCCTICAC GCAGGCIAIG ACCAGGIACT CCGCCCCCCC	8650
CGGGGACCCC CCACAACCAG AATACGACTT GGAGCTTATA ACATCATGCT	8700
CCTCCAACGT GTCAGTCGCC CACGACGCCG CTGGAAAGAG GGTCTACTAC	8750
CTTACCCGTG ACCCTACAAC CCCCCTCGCG AGAGCCGCGT GGGAGACAGC	8800
AAGACACACT CCAGTCAATT CCTGGCTAGG CAACATAATC ATGTTTGCCC	8850
CCACACTGTG GGCGAGGATG ATACTGATGA CCCATTTCTT TAGCGTCCTC	8900
ATACCACCE ATCACCITGA ACACCICIT AACIGICAGA TCTACCGACC	8950
CIGCIACICC ATAGAACCAC IGGATCIACC TOCAATCAIT CAAAGACICC	9000

FIG. 16E

ATGGCCTCAG	CGCATTTTCA	CTCCACAGTT	ACTOTOCAGG	TGAAATCAAT	9050
AGGGTGGCCG					9100
GAGACACCGG	GCCCGGAGCG	TOCGCCCTAG	CCTTCTGTCC	AGAGGAGGCA	9150
GGCTGCTAT	ATGIGGCAAG	TACCICTICA	ACTGGGCAGT	AAGAACAAAG	9200
CICAAACICA	CTCCAATAGC	GECCECTGEC	CCCCICGACT	TGICCGGITG	9250
GITCACGGCT	GGCTACAGCG	GGGGAGACAT	TTATCACAGC	GIGICICAIG	9300
CCCGGCCCCG	CIGGITCIGG	TITIGCCIAC	TOCTGCTOGC	TGCAGGGGTA	9350
CCCATCTACC	TOCTOCCCAA	CCGATGAAGG	TIGGGGTAAA	CACTCCGGCC	9400
TCTTAAGCCA	Triccigitt	TITTITITI	TTTTTTTT	TTTTTCTTT	9450
THEFT	TCCTTTCCTT	CITTITITICC	TITCITITIC	CCTTCTTTAA	9500
TEGTEGETICE	ATCTTAGCCC	TAGTCACGGC	TACCIGIGAA	AGGICCGIGA	9550
GCCGCATGAC	TGCAGAGAGT	GCTGATACTG	ecciciciec	AGATCATGT	9599

FIG. 16F

10 20 30 40 50	
1234567890 1234567890 1234567890 1234567890 1234567890	
MSINPKPORK IKRNINRRPO DVKFPGGGOI VGGVYLLPRR GPRLGVRAIR	50
KASERSOPRG RROPIPKARR PEGRAWAOPG YPWPLYGNEG LGWAGWLLSP	100
RGSRPSWGPT DPRRRSRNLG KVIDILICGF ADLMGYIPLV GAPLGGAARA	150
LAHGVRVLED GVNYATGNLP GCSFSIFILA LLSCLTIPAS AYEVRNVSGI	200
YHVINDOSNS SIVYEAADVI MHIPGOVPOV QEGNSSROW ALIPILAARN	250
ASVPTTTIRR HVDLLVGIAA FCSAMYVGDL CGSIFLVSQL FTFSPRRHET	300
VODCNOSIYP CHVSCHRMAW IMMMWSPIT ALVVSQLLRI PQAVVIMVAG	350
AHWGVLAGLA YYSMWGWAK VLIVALLFAG VDGETHTTGR VAGHTTSGFT	400
SLFSSGASQK IQLVNINGSW HINRIALNON DSLQIGFFAA LFYAHKFNSS	450
GCPERMASCR PIDWFAQGWG PITYIKPNSS DQRPYCWHYA PRPCGVVPAS	500
QVCGPVYCFT PSPVVVGTTD RSGVPTYSWG ENETDVMLLN NIRPPQGNWF	<b>55</b> 0
GCIMMISIGF TKICGGPPON IGGVGNRILI CPIDCFRKHP EATYTKOGSG	600
PWLTPRCLVD YPYRLWHYPC TLNFSIFKVR MYVGGVEHRL NAACIWIRGE	650
RCNLEDRDRS ELSPLLISTT EWQILPCAFT TLPALSTGLI HLHQNIVDVQ	700
YLYGVGSAFV SFAIKWEYIL LLFILLADAR VCACLWMMLL IAQAFAALEN	750
LVVINAASVA GAHGILSFLV FFCAAWYIKG RLAPGAAYAF YGWPLLLLL	800
LALPPRAYAL DTEVAASCGG VVLVGLMALT LSPYYKRYIS WCMWWLQYFL	850
TRVEAQLHWW VPPLNVRGGR DAVILLIMCVV HPTLVFDITK LLLAIFGPLW	900
ILOASILKVP YFVRVQGLLR ICALARKIAG GHYVQMAIIK LGALIGIYVY	950
NHLTPLRDWA HNGLRDLAVA VEPVVFSRME TKLTTWGADT AACGDIINGL	1000
PVSARROOFI LLGPADOMVS KOWRLLAPIT AYAQQIRGLL GCIITSLIGR	1050
DKNOVEGEVQ IVSTATQIFL ATCINGVCWT VYHGAGIRTI ASPKGPVIQM	1100
YTNVDQDLVG WPAPQGSRSL TPCTCGSSDL YLVTRHADVI PVRRRGDSRG	1150
SLLSPRPISY LKGSSGGPLL CPACHAVGLF RAAVCTRGVA KAVDFIPVEN	1200
LGTIMRSPVF TDNSSPPAVP QSFQVAHLHA PIGSGKSIKV PAAYAAQGYK	1250
VLVLNPSVAA TLGFGAYMSK AHGVDPNIRT GVRTITIGSP ITYSTYGKFL	1300
ADGGCSGGAY DILICDECHS TDATSLIGIG TVLDQAETAG ARLVVLATAT	1350
PPGSVIVSHP NIEEVALSIT GEIPFYGKAI PLEVIKGGRH LIFCHSKKKC	1400
DELAAKLVAL GINAVAYYRG LDVSVIPISG DVVVVSIDAL MIGFIGDFDS	1450
VIDONICVIQ TVDFSLDPIF TIETITLPQD AVSRTQRRGR TGRGKPGIYR	1500
FVAPGERPSG MFDSSVLCEC YDAGCAWYEL TPAETIVRLR AYMNTPGLPV	1550
CODHLEFWEG VFTGLTHIDA HFLSQTKQSG ENFPYLVAYQ ATVCARAQAP	1600
PPSWDOMWKC LIRLKPILHG PIPLLYRLGA VQNEVILIHP ITKYIMIOMS	1650
ADLEVVISTW VLVGGVLAAL AAYCLSTGCV VTVGRIVLSG KPAIIPDREV	1700
LYOFFDEMEE CSOHLPYTEQ CMMLAEQFKQ KALGLLQTAS RHAEVTTPAV	1750
OTMOKLEVF WAKHMINFIS GIQYLAGLST LPCNPAIASL MAFTAAVTSP	1800
LTTGOTLLFN ILGGWAAQL AAPGAATAFV GAGLAGAAIG SVGLGKVLVD	1850
ILAGYGAGVA GALVAFKINS GEVPSTEDLV NIIPAILSPG ALVVGVVCAA	1900

FIG. 16G

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### 40149

### H77CV-J4aa Sequence

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
TT PPHYGPGE	GAVOWMNRLI	AFASRGNHVS	PIHYVPESDA	AARVIAILSS	1950
TITATOLI RRI.	HOWISSECTT	PCSGSWLRDI	WDWICEVLSD	FKIWLKAKLM	2000
POT PETPEVS	CORGYRGWR	GDGIMHIRCH	CGAELIGHVK	NGIMRIVGPR	2050
TODAMAGGIF	PTNAYTIGEC	TPLPAPNYKF	ALWRVSAEEY	VEIRRVGDFH	2100
WINDSHITTINI.	KCPCOTPSPE	FFTELDGVRL	HRFAPPCKPL	LREEVSFRVG	2150
THEVERICOI.	PCEPEPDVAV	LISMLIDPSH	TTAFAAGRRL	ARGSPPSMAS	2200
CCACOT SAPS	TEXATOTANHO	SPDAELIEAN	LLWROEMGGN	TIRVESENKV	2250
לעון מכאומנו לע	AFFDEREVSV	PAETLRKSRR	FARALPWAR	<b>BDANBBLAET</b>	2300
	VHCCPI.PPPR	SPPVPPPRKK	RIVVLIESIL	STALAELATK	2350
CECCCOTOCT	TOTALITISSE	PAPSGCPPDS	DVESYSSMPP	LEGEPGDPDL	2400
CTCCMCTT/CC	CATTEDAVOC	SMSYSWIGAL	, VTPCAAEEQK	LPINALSNSL	2450
	TSRSACOROK	KVTFDRLQVL	, DSHYQDVLKE	VKAAASKVKA	2500
אוד ד כק זביציא ריכ	TIPPHSAKSK	FGYGAKDVRC	: HARKAVAHIN	SWKDLLELS	2550
T PORTECTION OF	KVIEVECVOPE	KGGRKPARLI	: VFPDLGVRVC	EKMALYL	2600
צד מן זק אוני	VGEOYSPGOE	VEFLVOAWKS	KKIPMLESYL	) TRUPLOIVIE	2650
CDIDIEEDI	Z COODI DECAE	R VAIKSLITERI	, YVGGPLINSF	( GEINGIRRUR	2700
		RACRAGLOI	CIMLVCGDDL	VVICESAGVQ	2750
EDAACI RAFT	r FAMTRYSAP!	GDPPOPEYDI	_ ELITSCSSN\	SVAHLGALKR	2800 ~
ידכרופייי דעעד	P PLARAMET	A RHTPVNSWLO	3 NIIMFAPILV	V ARMILMIHEE	2850
CAT TARDOL!	E OAT NOETYG	A CYSIEPLDLI	PIIQRLHGLS	5 AFSLHSYSPG	2900
ETNEKA ACT	R KT GTPPIRA	N RHRARSVRAJ	R LLSRGGRAA	L CGRATTHIMMAN	2950
דסיית דא דעירס	A AAGRIDISG	W FTAGYSGED	I YHSVSHARPI	R WEWECILLIA	3000
AGVGTYLLP					3011

FIG. 16H

# #1a. 3' Deletion mutants of pCV-H77C Sequence of 3' untranslated region of pCV-H77C

5'UTR ORF - 43 nts 81 nts poly U-UC 3' conserved 3' variable (3' variable region; 43 nts) region region TGA (Stop codon for polyprotein) AGGTTGGGGT AAACACTCCG GCCTCTTAAG CCATTTCCTG Afl II (poly U-UC region; 81 nts) CTTCTTTTT TCCTTTCTTT TTCCCTTCTT T (3' conserved region; 101 nts) AATGGTGGCT CCATCTTAGC CCTAGTCACG GCTAGCTGTG AAAGGTCCGT GAGCCGCATG ACTGCAGAGA GTGCTGATAC TGGCCTCTCT GCAGATCATG

#1a -1. pCV-H77C(-98X); 3' 98 nucleotides removed from pCV-H77C

TGAAGGTTGG GGTAAACACT CCGGCCTCTT AAGCCATTTC CTGTTTTTTT
TTTTTTTTTT TTTTTTTTTT TCTTTTTTT TTTCCTTTCCTTT
TTTTCCTTTC TTTTTCCCTT CTTTAAT

#1a -2. pCV-H77C(-42X); 3' 42 nucleotides removed from pCV-H77C

TGAAGGTTGG GGTAAACACT CCGGCCTCTT AAGCCATTTC CTGTTTTTT
TTTTTTTTTT TTTTTTTTTT TCTTTTTTT TTTCCTTTC
TTTTCCTTTC TTTTTCCCTT CTTTAATGGT GGCTCCATCT TAGCCCTAGT
CACGGCTAGC TGTGAAAGGT CCGTGAGCCG CAT

#1a -3. pCV-H77C(X-52); All of the 3' UTR sequence, except 3' 49 nucleotides, removed from pCV-H77C

TGAGCCGCAT GACTGCAGAG AGTGCTGATA CTGGCCTCTC TGCAGATCAT

FIG. 17A

#1a -4. pCV-H77C(X); All of the 3' UTR sequence, except 3' 101 nucleotides, removed from pCV-H77C

TGAAATGGTG GCTCCATCTT AGCCCTAGTC ACGGCTAGCT GTGAAAGGTC CGTGAGCCGC ATGACTGCAG AGAGTGCTGA TACTGGCCTC TCTGCAGATC ATGT

#1a -5. pCV-H77C(+49X); The proximal 49 nucleotides of the 3' conserved region ( 98 nucleotides; AAT not included) removed from pCV-H77C

#1a -6. pCV-H77C(VR-24); First 24 nucleotides of the 3' variable region removed from pCV-H77C

#1a -7. pCV-H77C(-U/UC); Poly U-UC region removed from pCV-H77C

TGAAGGTTGG GGTAAACACT CCGGCCTCTT AAGCCATTTC CTGAATGGTG

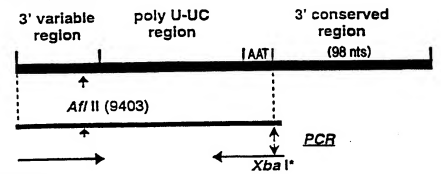
GCTCCATCTT AGCCCTAGTC ACGGCTAGCT GTGAAAGGTC CGTGAGCCGC

ATGACTGCAG AGAGTGCTGA TACTGGCCTC TCTGCAGATC ATGT

**FIG. 17B** 

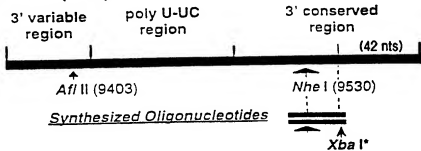
### #1b. Strategy of 3' Deletion mutants

#### #1b -1. pCV-H77C(-98X)



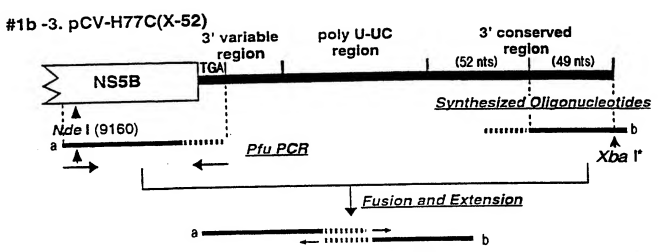
- 1. PCR Amplification
- 2. Purification of PCR products
- 3. Digestion with Afl II and Xba I
- 4. Cloning of Afl II IXba I fragment into pCV-H77C
- 5. Complete sequence analysis
- 6. in vitro transcription (within 24 hours of inoculation)
- 7. Percutaneous intrahepatic transfection into chimpanzee; 11/26/97 and 12/17/97
- 8. Result : Negative ( No replication)

#### #1b -2. pCV-H77C(-42X)



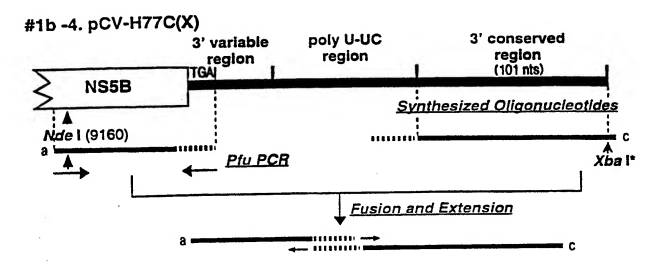
- 1. Synthesis of oligonucleotides (sense and anti-sense)
- 2. Hybridization of oligonucleotides
- 3. Digestion with Nhe I and Xba I
- 4. Cloning of Nhe I /Xba I fragment into pG9-KL26 (3' UTR of H77C)
- 5. Sequence analysis
- 6. Cloning of 3' UTR (-42X) [Afl II /Xba I fragment] into pCV-H77C
- 7. Complete sequence analysis
- 8. in vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee (Schedule; 1/22/98, 2/5/98)

### **FIG. 17C**



- 1. Fragment a; Pfu PCR amplification and purification
- 2. Fragment b; Synthesized oligonucleotides (anti-sense)
- 3. Fusion and extension
- 4. TA cloning
- Sequence analysis
- 6. Cloning Nde I-Xba I fragment with correct sequence into pCV-H77C
- 7. Complete sequence analysis
- 8. In vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee

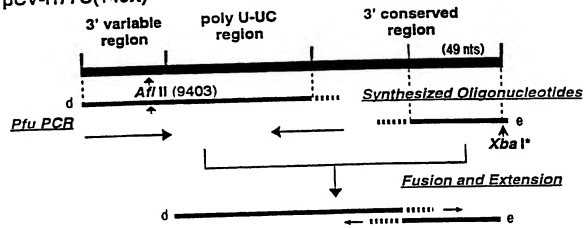
FIG. 17D



- 1. Fragment a; Pfu PCR amplification and purification
- 2. Fragment c; Synthesized oligonucleotides (anti-sense)
- 3. Fusion and extension
- 4. TA cloning
- 5. Sequence analysis
- 6. Cloning Nde I-Xba I fragment with correct sequence into pCV-H77C
- 7. Complete sequence analysis
- 8. In vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee

**FIG. 17E** 

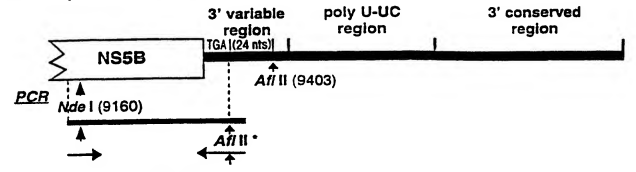
#1b -5. pCV-H77C(+49X)



- 1. Fragment d; Pfu PCR amplification and purification
- 2. Fragment e ; Synthesized oligonucleotides (anti-sense)
- 3. Fusion and extension
- 4. TA cloning
- 5. Sequence analysis
- 6. Cloning Aff II-Xba I fragment with correct sequence into pCV-H77C
- 7. Complete sequence analysis
- 8. In vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee

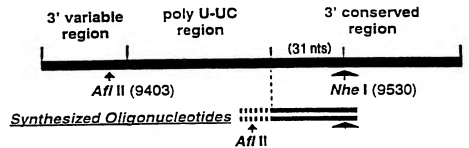
FIG. 17F

#### #1b -6. pCV-H77C(VR-24)



- 1. PCR Amplification
- 2. Purification of PCR products
- 3. Digestion with Nde I and Afl I
- 4. Cloning of Nde I /Afl II fragment into pCV-H77C
- 5. Complete sequence analysis
- 6. in vitro transcription (within 24 hours of inoculation)
- 7. Percutaneous intrahepatic transfection into chimpanzee

#### #1b -7. pCV-H77C(-U/UC)



- 1. Synthesis of oligonucleotides ( sense and anti-sense )
- 2. Hybridization of oligonucleotides
- 3. Digestion with Aff II and Nhe I
- 4. Cloning of Afl II and Nhe I fragment into pG9-KL26
- 5. Sequence analysis
- 6. Cloning of 3' UTR ( -poly U-UC ) [Afl II /Xba I fragment] into pCV-H77C
- 7. Complete sequence analysis
- 8. in vitro transcription (within 24 hours of inoculation)
- 9. Percutaneous intrahepatic transfection into chimpanzee

### **FIG. 17G**

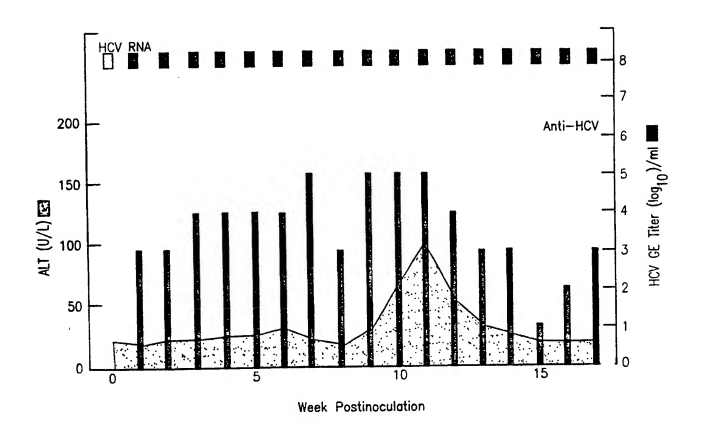


FIG. 18A

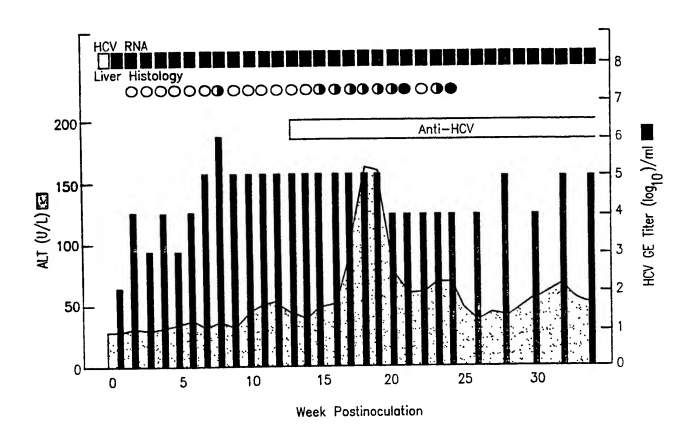


FIG. 18B

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#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) Internati nal Patent Classificati n 6:

C12N 15/40, C07K 14/18, C12Q 1/70, C07K 16/10, A61K 39/29

A3

(11) International Publication Number:

WO 99/04008

(43) International Publication Date:

28 January 1999 (28.01.99)

(21) International Application Number:

PCT/US98/14688

(22) International Filing Date:

16 July 1998 (16.07.98)

(30) Priority Data:

60/053,062 09/014,416

18 July 1997 (18.07.97) US 27 January 1998 (27.01.98)

US

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

With international search report.

(88) Date of publication of the international search report: 1 April 1999 (01.04.99)

(54) Title: CLONED GENOMES OF INFECTIOUS HEPATITIS C VIRUSES AND USES THEREOF

#### (57) Abstract

The present invention discloses nucleic acid sequences which encode infectious hepatitis C viruses and the use of these sequences, and polypeptides encoded by all or part of these sequences, in the development of vaccines and diagnostics for HCV and in the development of screening assays for the identification of antiviral agents for HCV.

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CLASSIFICATION OF SUBJECT MATTER C 6 C12N15/40 C07k A61K39/29 IPC 6 C07K16/10 C07K14/18 C12Q1/70According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C12N C07K C12Q A61K IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category 5 Citation of document, with indication, where appropriate, of the relevant passages YOO B J ET AL: "Transfection of a 1,13-24. X 33 - 35differentiated human hepatoma cell line 42.43 (Huh7) with in vitro-transcribed hepatitis C virus (HCV) RNA and establishment of a long-term culture persistently infected with HCV" JOURNAL OF VIROLOGY, vol. 69, no. 1, January 1995, pages 32-38, XP002022696 AMERICAN SOCIETY FOR MICROBIOLOGY US see the whole document Patent family members are listed in annex. Further documents are listed in the continuation of box C. lχ Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 22/01/1999 12 January 1999 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Cupido, M

1

Internatic Application No
PCT/US 98/14688

		PC1/US 98/14688			
C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT					
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.			
E	WO 98 39031 A (UNIVERSITY OF WASHINGTON; KOLYKHALOV A; RICE C) 11 September 1998  see SEQ ID NO:1, representing a HCV H77 consensus sequence having 99.6% identity with the nucleic acid sequences in figures 4A-4F of the present application.	1,4,5, 13-24, 33-43			
X	EP 0 516 270 A (JAPAN IMMUNO INC) 2 December 1992 see the whole document	23,24, 40,41			
Α	WO 97 08310 A (WASHINGTON UNIVERSITY) 6 March 1997 see page 15; figure 3	1-43			
P, X	YANAGI M ET AL: "Transcripts of a chimeric cDNA clone of hepatitis C virus genotype 1b are infectious in vivo" VIROLOGY, vol. 244, no. 1, 25 April 1998, pages 161-172, XPO2089701 ORLANDO US see the whole document	1,6-10, 13-24, 33-43			

Intern unal application No.

PCT/US 98/14688

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:  1. X Claims Nos.:	Box I Observations wher certain claims were found unsearchable (Continuation of item 1 of first sheet)
because they relate to subject matter not required to be searched by this Authority, namely:  Remark: Although claim(s) 41 and 43  is(are) directed to a method of treatment of the animal body, the search has been carried out and based on the alleged effects of the composition.  2. X Claims Nos.: 29 and 30 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:  The scope of claims 29 and 30 is so unclear and not well specified that a meaningful search was not possible  3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).  B x II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)  This International Searching Authority found multiple inventions in this international application, as follows:  1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.  2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.  3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report	This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
a meaningful search was not possible  3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule6.4(a).  B x II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)  This International Searching Authority found multiple inventions in this international application, as follows:  1. As all required additional search fees were timely paid by the applicant, this international Search Report covers all searchable claims.  2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.  3. As only some of the required additional search fees were timely paid by the applicant, this international Search Report	because they relate to subject matter not required to be searched by this Authority, namely:  Remark: Although claim(s) 41 and 43  is(are) directed to a method of treatment of the animal body, the search has been carried out and based on the alleged effects of the composition.  2. X Claims Nos.: 29 and 30 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).  B x II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)  This International Searching Authority found multiple inventions in this international application, as follows:  1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.  2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.  3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report	a meaningful search was not possible
This International Searching Authority found multiple inventions in this international application, as follows:  1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.  2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.  3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report	
<ol> <li>As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.</li> <li>As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.</li> <li>As only some of the required additional search fees were timely paid by the applicant, this International Search Report</li> </ol>	B x II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
<ul> <li>searchable claims.</li> <li>As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.</li> <li>As only some of the required additional search fees were timely paid by the applicant, this International Search Report</li> </ul>	This International Searching Authority found multiple inventions in this international application, as follows:
of any additional fee.  3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest  The additional search fees wer accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.	

Information on patent family members

Internatio	pplication No
PCT/US	98/14688

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